

Trust-Region Based Return Mapping Algorithm for Implicit Integration of Elastic-Plastic Constitutive Models

B. T. Lester* and W. M. Scherzinger

Solid Mechanics Department, Sandia National Laboratories, Albuquerque, NM 87185, USA

SUMMARY

A new method for the solution of the non-linear equations forming the core of constitutive model integration is proposed. Specifically, the trust-region method that has been developed in the numerical optimization community is successfully modified for use in implicit integration of elastic-plastic models. Although attention here is restricted to these rate-independent formulations, the proposed approach holds substantial promise for adoption with models incorporating complex physics, multiple inelastic mechanisms, and/or multiphysics. As a first step, the non-quadratic Hosford yield surface is used as a representative case to investigate computationally challenging constitutive models. The theory and implementation are presented, discussed, and compared to other common integration schemes. Multiple boundary value problems are studied and used to verify the proposed algorithm and demonstrate the capabilities of this approach over more common methodologies. Robustness and speed are then investigated and compared to existing algorithms. Through these efforts, it is shown that the utilization of a trust-region approach leads to superior performance versus a traditional closest-point projection Newton-Raphson method and comparable speed and robustness to a line search augmented scheme. Copyright © 0000 John Wiley & Sons, Ltd.

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KEY WORDS: Constitutive Model Integration; Return Mapping Algorithms; Trust-Region; Hosford Yield Surface

1. INTRODUCTION

As the phenomenological theory of plasticity has matured in recent decades, the models being proposed have grown increasingly complex. For instance, non-quadratic yield surfaces like those put forth by Hosford [1], the classical anisotropic expression of Hill [2], and more recent models that are both non-quadratic and anisotropic (*i.e.* Barlat and coworkers [3, 4], Karafillis and Boyce [5], and Cazacu and coworkers [6, 7]) have all been developed and implemented. The additional capabilities provided by such models have been useful in the analysis of sheet metal forming [8, 9, 10] and inflation/burst of tubes [11, 12, 13] in which accurately capturing the anisotropic plastic flow is essential.

Similar to the aforementioned plasticity models, structural analyses and corresponding simulations have also grown in terms of complexity and size. In turn, there is an increasing focus on the efficient implicit integration of both the global finite element problem and local stress updating procedure. With respect to the latter problem, the fully implicit closest point projection (CPP) and semi-implicit convex cutting plane (CCP) return mapping algorithms (RMAs) popularized by Simo, Ortiz, and coworkers [14, 15, 16, 17, 18, 19] have been extensively explored. Such implementations have been pursued for a wide variety of phenomenological constitutive models incorporating different physics and flow rules (*e.g.* [20, 21, 22, 23, 24, 25]). A modified approach

*Correspondence to: btleste@sandia.gov

has been taken by Yoon *et al.* [26, 27, 8], who have proposed a return mapping scheme based on the control of a potential residual. In all of these cases, the Newton-Raphson method is used to solve the non-linear equations forming the core of the stress updating approaches.

Unlike the radial return approaches common with J_2 plasticity, pure Newton-Raphson solvers are not guaranteed to converge for more complex models. Comprehensive studies by Pérez-Foguet and Armero [28] and Scherzinger [29] have extensively explored this issue for a variety of models and demonstrated the insufficiency of a Newton-Raphson solver under many conditions. For implicit global solvers, the ability of the constitutive routine to converge under any input deformation is important to avoid costly cutbacks to the global timestep. This is especially true for non-linear finite element codes like Sierra/SM [30] that utilize an iterative global solver and therefore may see a wider range of trial deformations during the solution process.

To improve the convergence properties, a variety of modifications to the underlying algorithms have been proposed. One of the more common approaches is to break up large input deformations into a series of smaller loadings and sequentially solve the subloadings until the intended total deformation is achieved. Conceptually, such schemes are similar to the arc-length methods [31] that are used with global FE solvers in which a constraint equation is solved to restrict load (and/or displacement) increments thereby improving robustness. Although more common with explicit stress integration schemes, these substepping approaches have been adopted by Yoon and colleagues [26, 27, 8], Scifert *et al.* [32, 33], and Rabahallah and coworkers [34] for a wide variety of constitutive models. Such methods, however, mitigate some of the advantages of a fully implicit scheme and introduce additional complexity to the ensuing implementation. An alternative to this has been the adoption of more complex numerical schemes to ensure convergence – notably line search methods. Early works like those of Dutko *et al.* [35] observed improvements in adding an additional line search step while more recent efforts such as those by Seifert and coworkers [36], Pérez-Foguet and Armero [28], and Scherzinger [29] have all extensively explored and demonstrated these benefits. For instance, Scherzinger noted convergence of the line search augmented CPP method for any trial stresses whose effective measure was less than or equal to 30 times that of yield for both the Hosford and Barlat models – far exceeding the performance of a pure Newton-Raphson solver.

These previous efforts (substepping and line search schemes) have focused on modifying or augmenting the Newton-Raphson solver to improve robustness. Given the developments in non-linear optimization and numerical methods since the early efforts of Simo, Ortiz, Hughes, and others investigating RMAs, an alternative approach to consider is the utilization of a new solver scheme with the CPP-RMA problem. One such possibility are *trust-region* based methods [37]. At their core, these algorithms search for updated solution variables minimizing an objective function within a trusted solution variable domain. Although primarily studied for general optimization problems [38, 39, 40, 41, 42], trust-region methods have been successfully adopted in structural design optimization [43] and flow control [44] applications. A previous effort by Shterenlikht and Alexander [45] attempted to use open-source implementations of both Levenberg-Marquardt and dogleg trust-region methods to integrate the Gurson-Tvergaard-Needleman (GTN) model. In their investigation, the former approach showed promise while severe issues with scaling were encountered with the latter leading to substantial convergence difficulty.

In this work, the possibility of using a tailored trust-region approach for the CPP-RMA problem to integrate elastic-plastic constitutive models is explored. Specifically, the numerical formulation of a Hosford plasticity model using a trust-region solution method is developed and implemented. A series of boundary value problems are then investigated to demonstrate the capabilities of this novel methodology and investigate the performance characteristics (especially convergence) in comparison to established Newton-Raphson and line search augmented approaches. To this end, this work is organized as follows. Section 2 describes the elastic-plastic model and its numerical implementation. Both the theoretical formulation and some basic discussion of the three solution methods – Newton-Raphson (NR), line search augmented Newton-Raphson (LS-NR), and the trust-region approach (TR) – are presented along with details of the trust-region implementation.

Numerical results and discussion are presented in Section 3 while concluding remarks are given in Section 4.

2. TRUST-REGION IMPLEMENTATION OF ELASTIC-PLASTIC MODELS

To investigate trust-region based constitutive model integration, a hypoelastic implementation of the Hosford model [1] is developed. This selection is made as it is a relatively simple form (only two parameters and isotropic) while still being able to induce substantial curvature in the yield surface. It is noted that the algorithm is presented in a sufficiently general fashion so that it is easily adaptable for use with other yield surfaces or isotropic hardening laws[†]. The theoretical model is presented in the next section while the numerical implementation is presented in Section 2.2.

2.1. Hosford Plasticity Model

Like many other rate-independent elastic-plastic models, an additive split in the strain tensor is assumed such that,

$$\varepsilon_{ij} = \varepsilon_{ij}^e + \varepsilon_{ij}^p \quad (1)$$

where ε_{ij} , ε_{ij}^e , and ε_{ij}^p are the total, elastic, and plastic strain tensors, respectively. The symmetric Cauchy stress, σ_{ij} , is given as,

$$\sigma_{ij} = \mathbb{C}_{ijkl} \varepsilon_{kl}^e = \mathbb{C}_{ijkl} (\varepsilon_{kl} - \varepsilon_{kl}^p), \quad (2)$$

with \mathbb{C}_{ijkl} being the fourth order elastic stiffness tensor (assumed isotropic). This leads to a stress increment equation[‡] of the form,

$$d\sigma_{ij} = \mathbb{C}_{ijkl} (d\varepsilon_{kl} - d\varepsilon_{kl}^p). \quad (3)$$

Assuming an associative flow rule the plastic strain increment, $d\varepsilon_{ij}^p$, can be written as

$$d\varepsilon_{ij}^p = d\gamma \frac{\partial f}{\partial \sigma_{ij}}, \quad (4)$$

where $d\gamma$ is the plastic consistency multiplier and f is the yield function describing the domain of elastic deformation. In this case, because of the associated flow rule the consistency multiplier is equal to effective plastic strain increment ($d\gamma = d\bar{\varepsilon}^p$). The plastic multiplier is found through consistency relations resulting from the traditional Kuhn-Tucker conditions,

$$d\gamma \geq 0; \quad d\gamma f = 0; \quad f \leq 0. \quad (5)$$

Bearing this in mind, the response of the model is largely dictated by the yield surface and hardening rule considered. In this case, the yield function is given by

$$f(\sigma_{ij}, \bar{\varepsilon}^p) = \phi(\sigma_{ij}) - \sigma_y(\bar{\varepsilon}^p) \quad (6)$$

where σ_y is the yield stress, $\bar{\varepsilon}^p$ is the isotropic hardening variable (equivalent plastic strain) and ϕ is the Hosford effective stress [1] defined as

$$\phi(\sigma_{ij}) = \left[\frac{|\sigma_1 - \sigma_2|^a + |\sigma_2 - \sigma_3|^a + |\sigma_1 - \sigma_3|^a}{2} \right]^{1/a}, \quad (7)$$

[†]Kinematic hardening is not treated in this work although the authors see no reason that would preclude an extension to such models.

[‡]As the focus of this effort is on numerical methods for solving local constitutive relations, discussions towards objective stress rates for hypoelastic formulations are neglected. For details and discussions of such issues, the reader is referred to [46, 47, 23, 32].

with σ_i being the principal stresses and a the fitting exponent ($1 \leq a \leq \infty$) giving the model its non-quadratic nature. In the developed algorithm, the numerical routine put forth by Scherzinger and Dohrmann [48] is used to determine the principal stresses. Linear hardening of the form,

$$\sigma_y(\bar{\varepsilon}^p) = \sigma_y^0 + K\bar{\varepsilon}^p, \quad (8)$$

is considered in which σ_y^0 is a constant initial yield stress and K the hardening modulus. Additionally, for this study it is assumed that the loadings are isothermal and temperature dependence is neglected. Details of this model, notably the derivatives of the effective stress, may be found in a previous effort [29].

2.2. Numerical Implementation

Regardless of numerical solution scheme (NR, LS-NR, or TR), the underlying non-linear stress updating problem is the same. Specifically, to numerically integrate the hypoelastic constitutive response posed in Section 2.1, an elastic predictor-inelastic corrector scheme based on operator splitting [14] is adopted. By assuming the given loading increment does not lead to inelastic deformation, an elastic trial stress, σ_{ij}^{tr} , is computed as,

$$\sigma_{ij}^{\text{tr}} = \sigma_{ij}^{(n)} + \mathbb{C}_{ijkl} d\varepsilon_{kl}^{(n+1)} \quad (9)$$

with the superscripts “ (n) ” and “ $(n+1)$ ” referring to variables at those load steps and $d\varepsilon_{ij}$ is the total strain increment. The validity of the elastic step may be determined by computing $f^{\text{tr}} = f(\sigma_{ij}^{\text{tr}}, \bar{\varepsilon}^p(n))$. If $f^{\text{tr}} < 0$, the elastic step is valid and the state variables are updated ($\sigma_{ij}^{(n+1)} = \sigma_{ij}^{\text{tr}}$, $\bar{\varepsilon}^p(n+1) = \bar{\varepsilon}^p(n)$). The more interesting plastic loading case occurs when $f^{\text{tr}} > 0$.

The solution to the inelastic correction is associated with the constrained optimization problem corresponding to maximum dissipation – see [19, 49] for discussions on the thermodynamics. During the inelastic correction, the solution is found by iteratively refining the state variables until a thermodynamically admissible state is achieved. For implicit methods like the CPP approach pursued here, this process is performed by enforcing the consistency relation and the plastic strain flow rule on the material state at $t = t_{n+1}$. Mathematically, these two conditions are given in residual form as

$$\{\mathbf{r}(\{\mathbf{x}\})\}^{(n+1)} = \begin{Bmatrix} r_{ij}^{\varepsilon(n+1)} \\ r^f(n+1) \end{Bmatrix} = \begin{Bmatrix} -d\varepsilon_{ij}^{p(n+1)} + d\gamma^{(n+1)} \frac{\partial \phi}{\partial \sigma_{ij}^{(n+1)}} \\ f(\sigma_{ij}^{(n+1)}, d\gamma^{(n+1)}) \end{Bmatrix} = \begin{Bmatrix} 0 \\ 0 \end{Bmatrix}, \quad (10)$$

where $\{\mathbf{x}\}$ are the solution variables that may be written as,

$$\{\mathbf{x}\}^{(n+1)} = \begin{bmatrix} \sigma_{ij}^{(n+1)} \\ d\gamma^{(n+1)} \end{bmatrix}. \quad (11)$$

The first, r_{ij}^{ε} , and second, r^f , residuals are associated with the flow rule and consistency condition, respectively.[§]

The non-linear problem posed by $\{\mathbf{r}\} = \{\mathbf{0}\}$ is solved numerically by iterating until convergence has been achieved. In this work, a merit function of the form,

$$\psi(\{\mathbf{x}\}) = \frac{1}{2} \{\mathbf{r}(\{\mathbf{x}\})\}^T [\mathbf{D}^1]^T [\mathbf{D}^1] \{\mathbf{r}(\{\mathbf{x}\})\} \quad (12)$$

[§]Given the mixing of scalar and tensorial variables in the residual and corresponding terms, in what follows a matrix notation will be used to compactly write and combine the various equations. Square brackets “[.]” will be used for matrix-like quantities while braces “{.}” shall be reserved for columnar vectors. The components of these different representations may be either tensorial or scalar and bold-face text is used to distinguish these objects.

in which $[\mathbf{D}^1]$ is a constant, diagonal matrix taking the form,

$$[\mathbf{D}^1] = \begin{bmatrix} c^e \mathbb{I}_{ijkl} & 0_{ij} \\ 0_{ij} & c^f \end{bmatrix}, \quad (13)$$

where \mathbb{I}_{ijkl} is the fourth order identity tensor, is used to assess convergence. Specifically, the square root of the merit function ($\sqrt{\psi}$) is used as the convergence criterion. Essentially, these constants are introduced as weights to appropriately scale the contributions of the different components of the residual. From Eqn. 10 it is noted that in the residual and solution variable vectors, both strain and stress quantities appear. Depending on the units being used, the difference in the values of the variables can be many orders of magnitude, highlighting the necessity of appropriately scaling the problem. Importantly, regardless of the values of c^e and c^f the minimum is achieved at $r_{ij}^e = r^f = 0$. By setting the constants to be one, $[\mathbf{D}^1]$ reduces to the identity matrix and the more traditional merit function of the form $\psi = (1/2) \{\mathbf{r}\}^T \{\mathbf{r}\}$ is recovered. Trust-region methods, however, are more susceptible to scaling issues as experienced by Shterenlikht and Alexander [45]. In the remainder of this work, an equal weight, stress-normalization is introduced by setting $c^e = E/\sigma_y^0$ and $c^f = 1/\sigma_y^0$ such that the merit function may be written as,

$$\psi(\{\mathbf{r}\}) = \frac{1}{2} \left(\left(\frac{E}{\sigma_y^0} \right)^2 r_{ij}^e r_{ij}^e + \left(\frac{r^f}{\sigma_y^0} \right)^2 \right). \quad (14)$$

The impact of this scaling on the algorithm performance is explored in Section 3.3.

To begin the inelastic correction process, the state variable values at the $k = 0$ iteration are initialized to those of the trial state while $d\varepsilon_{ij}^{p(k=0)} = 0$ and $d\gamma^{(k=0)} = 0$.[¶] For all three numerical solution schemes considered here, the solution variables are iteratively updated via

$$\{\mathbf{x}\}^{(k+1)} = \{\mathbf{x}\}^{(k)} + \alpha^{(k)} \{\mathbf{p}\}^{(k)}, \quad (15)$$

where $\alpha^{(k)}$ and $\{\mathbf{p}\}^{(k)}$ are the step size and vector, respectively, and $\{\mathbf{p}\} = [p_{ij}^\sigma, p^\gamma]^T$ with $p_{ij}^\sigma = \Delta\sigma_{ij}$ and $p^\gamma = \Delta\gamma$. The three approaches (NR, LS-NR, and TR) differ in how the step size, $\alpha^{(k)}$, and step vector, $\{\mathbf{p}\}^{(k)}$, are determined.

In the simplest and most-common approach (NR), the step size is fixed ($\alpha^{(k)} = 1$) and the step vector, $\{\mathbf{p}^{\text{NR}}\}^{(k)}$, takes the classical form of,

$$\{\mathbf{p}^{\text{NR}}\}^{(k)} = -[\mathbf{J}]_{(k)}^{-1} \{\mathbf{r}\}^{(k)} \quad (16)$$

with $[\mathbf{J}]$ being the Jacobian defined as $[\mathbf{J}] = \frac{\partial\{\mathbf{r}\}}{\partial\{\mathbf{x}\}}$. Given the residuals in Eqn. 10, the Jacobian is

$$[\mathbf{J}] = \begin{bmatrix} (\mathcal{L}_{ijkl})^{-1} & \frac{\partial\phi}{\partial\sigma_{ij}} \\ \frac{\partial\phi}{\partial\sigma_{ij}} & -\frac{\partial\phi}{\partial\varepsilon^p} \end{bmatrix}, \quad (17)$$

where \mathcal{L}_{ijkl} is given by,

$$\mathcal{L}_{ijkl} = \left(\mathbb{C}_{ijkl}^{-1} + d\gamma \frac{\partial^2\phi}{\partial\sigma_{ij}\partial\sigma_{kl}} \right)^{-1}. \quad (18)$$

Line search approaches vary from NR and other methods by selecting a step direction (in this case, the NR direction) and then finding a step size minimizing the merit function in the domain $\alpha^{(k)} = (0, 1]$. This constraint serves to enforce that each iteration reduces the merit function. As

[¶]In the remainder of this section, it shall be assumed that unless specifically noted the variable values correspond to those at the “(n + 1)” timestep and “(k)” and “(k + 1)” superscripts are to denote the previous and current correction iterations. Furthermore, the term increment shall be reserved for use with the global time/loadstep while iteration shall be used to describe the inelastic correction process.

an exact minimum can be expensive to compute, an approximate minimum is often found by a quadratic approximation. In practice, the utilization of a line search approach is fairly straightforward as it only requires an additional step to find $\alpha^{(k)}$. For details of the implemented LS-NR scheme for the Hosford model, see [29].

Unlike line search methods that directly investigate the merit function, trust-region based schemes instead construct a local approximation to the scalar function to be minimized via a model problem, $m^{(k)}(\{\mathbf{p}\})$, and find the step vector that minimizes the model problem. As this representation is only an approximation of the actual non-linear problem, the step vector is only found in the solution variable space sufficiently close to the current solution that can be trusted (hence the name). This domain is defined to be the region in solution variable space within a ball of radius $\Delta^{(k)}$ ($\|\{\mathbf{p}\}^{(k)}\| \leq \Delta^{(k)}$). The size of this trust-region is then updated depending on the improvement (or lack thereof) of a step vector. Limiting the solution space serves to highlight a key distinction between line search and trust-region methods. Namely, the former picks a step vector and then determines the size while the latter selects the step size and then finds the step vector. [37]

In the current study, a quadratic approximation of the form [37],

$$m^{(k)}(\{\mathbf{p}\}) = \Psi^{(k)} + \{\mathbf{g}\}_{(k)}^T \{\mathbf{p}\} + \frac{1}{2} \{\mathbf{p}\}^T [\mathbf{B}]^{(k)} \{\mathbf{p}\}, \quad (19)$$

is used for the model problem where $\Psi^{(k)}$ is the function to be minimized, $\{\mathbf{g}\}^{(k)}$ is the gradient of $\Psi^{(k)}$ ($\{\nabla \Psi\}^{(k)}$), and $[\mathbf{B}]^{(k)}$ is an approximation of the Hessian ($[\nabla^2 \Psi]^{(k)}$). When $\Psi^{(k)}$ is an unweighted form of Eqn. 12 ($c^e = c^f = 1$), these terms are commonly written as,

$$\Psi^{(k)} = \frac{1}{2} \{\mathbf{r}\}_{(k)}^T \{\mathbf{r}\}^{(k)}; \quad \{\mathbf{g}\}^{(k)} = [\mathbf{J}]_{(k)}^T \{\mathbf{r}\}^{(k)}; \quad [\mathbf{B}]^{(k)} = [\mathbf{J}]_{(k)}^T [\mathbf{J}]^{(k)}. \quad (20)$$

Schematically, this method is illustrated in Fig. 1 in which a simplified case of J_2 plasticity with no hardening is treated. In this case, contours corresponding to a representative model problem, $m^{(k)}$, (whose chosen form is selected for demonstration purposes) are presented alongside the yield surface and initial and final stress states. Two circles of radii Δ^1 and Δ^2 corresponding to different domains along with their minimization vectors, $\{\mathbf{p}^1\}$ and $\{\mathbf{p}^2\}$, are also presented. In this visualization, the contributions of the effective plastic strain increment, $d\gamma^{(k)}$, on the trust-region size, $\Delta^{(k)}$, are neglected for clarity of presentation. Note, in this case the two step vectors $\{\mathbf{p}^1\}$ and $\{\mathbf{p}^2\}$ do not represent sequential correction step vectors but are instead two different solutions arising from distinct TR radii. Importantly, as is observed from Fig. 1, it is noted that changing the trust-region radius from Δ^1 to Δ^2 not only changes the magnitude of the step direction vectors ($\{\mathbf{p}^1\}$ and $\{\mathbf{p}^2\}$) but also the direction highlighting the impact of this parameter and flexibility of the method. Successive iterations adjust the size of the trust-region based on measures of improvement and algorithmic rules. A corresponding refined model problem is also produced that is then minimized over the next iteration. Through this iterative process, the material state converges on the correct solution.

As mentioned earlier, variable scaling may substantially affect convergence for the non-linear problem corresponding to constitutive model integration and needs to be handled appropriately. One common way to deal with this is to introduce the scaling ahead of time and develop a modified model problem, $\tilde{m}^{(k)}$, that is of the general form given in Eqn. 19 [37]. In this way, the contributions of the different solution variables equally contribute and all of the developments towards solving the problems posed by Eqns. 19 and 20 can be leveraged.

To come up with the scaled model problem, $\tilde{m}^{(k)}$, two areas need to be addressed. The first has already been discussed in this section – the merit function. Specifically, the merit function serves as the function to be minimized such that $\psi^{(k)}$ plays the role of $\Psi^{(k)}$ in the scaled problem. The second scaling issue that needs to be addressed is associated with the differences in the solution variables and is manifest in the step vector, $\{\mathbf{p}\}^{(k)}$. A similar approach to that of the merit function is used and a scaled step vector, $\{\tilde{\mathbf{p}}\}^{(k)}$, is introduced such that,

$$\{\tilde{\mathbf{p}}\}^{(k)} = [\mathbf{D}^2] \{\mathbf{p}\}^{(k)}, \quad (21)$$

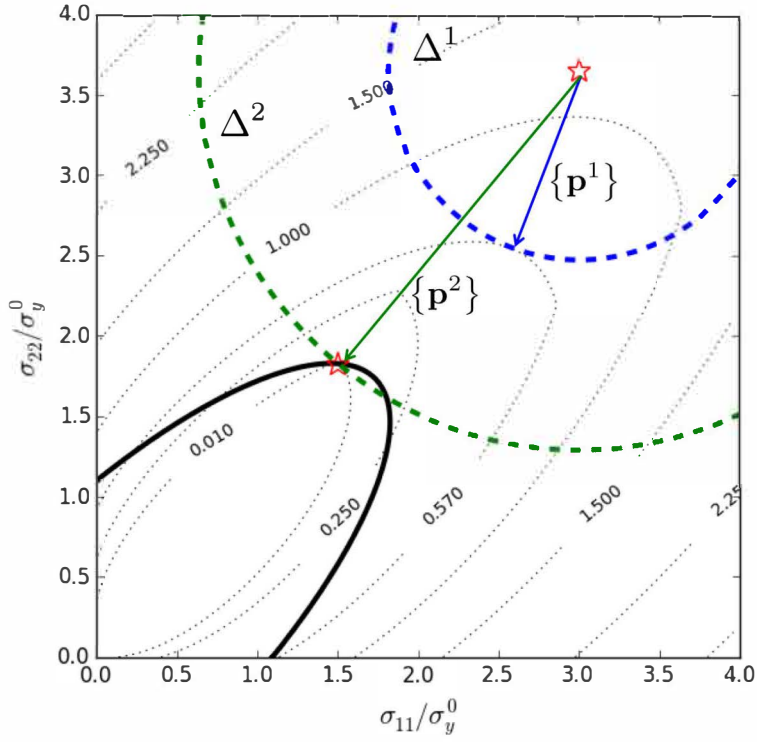


Figure 1. Schematic representation of the TR correction approach. A J_2 yield surface (solid black line) is marked along with an initial and final stress state (red stars). Dotted black lines correspond to the contours of a representative model problem, $m^{(k)}$. Two different trust-region domains (and their radii, $\Delta^{1,2}$) are marked

along with the corresponding step direction vector, $\{p^{1,2}\}$.

with $[D^2]$ taking the form,

$$[D^2] = \begin{bmatrix} b^\sigma \mathbb{I}_{ijkl} & 0_{ij} \\ 0_{ij} & b^\gamma \end{bmatrix}. \quad (22)$$

A scaled trust-region magnitude, $\tilde{\Delta}^{(k)}$, is also introduced such that $\|\{\tilde{p}\}^{(k)}\| \leq \tilde{\Delta}^{(k)}$. For these scalings, a stress normalization is again used such that $b^\sigma = 1$ and $b^\gamma = 2\mu$. Using $\psi^{(k)}$ as the merit function and Eqn. 21 to write the scaled model problem, $\tilde{m}^{(k)}$, in terms of the scaled step vector produces an expression of the form,

$$\tilde{m}^{(k)}(\{\tilde{p}\}) = \psi^{(k)} + \{\tilde{g}\}_{(k)}^T \{\tilde{p}\} + \frac{1}{2} \{\tilde{p}\}^T [\tilde{B}]^{(k)} \{\tilde{p}\}, \quad (23)$$

where the introduction of $\{\tilde{p}\}$ leads to the following relation for the gradient,

$$\{\tilde{g}\}^{(k)} = [D^2]^{-T} \left([D^1] [J]^{(k)} \right)^T [D^1] \{r\}^{(k)}, \quad (24)$$

and Hessian,

$$[\tilde{B}]^{(k)} = [D^2]^{(-T)} \left([D^1] [J]^{(k)} \right)^T \left([D^1] [J]^{(k)} \right) [D^2]^{(-1)}. \quad (25)$$

More convenient forms of these expressions may be written,

$$\{\tilde{\mathbf{g}}\}^{(k)} = \left[\begin{array}{c} \frac{(c^e)^2}{b^\sigma} r_{kl}^{\varepsilon(k)} \mathcal{L}_{kl}^{-1} + \frac{(c^f)^2}{b^\sigma} r_{ij}^{f(k)} \frac{\partial \phi}{\partial \sigma_{ij}^{(k)}}, \quad \frac{(c^e)^2}{b^\gamma} r_{ij}^{\varepsilon(k)} \frac{\partial \phi}{\partial \sigma_{ij}^{(k)}} - \frac{(c^f)^2}{b^\gamma} r_{ij}^{f(k)} \frac{\partial \sigma_y}{\partial \varepsilon^{p(k)}} \end{array} \right]^T, \quad (26)$$

and,

$$[\tilde{\mathbf{B}}]^{(k)} = \left[\begin{array}{cc} \left(\frac{c^e}{b^\sigma} \right)^2 \mathcal{L}_{ijrs}^{-1} \mathcal{L}_{rskl}^{-1} + \left(\frac{c^f}{b^\sigma} \right)^2 \frac{\partial \phi}{\partial \sigma_{ij}^{(k)}} \frac{\partial \phi}{\partial \sigma_{kl}^{(k)}}, & \frac{(c^e)^2}{b^\sigma b^\gamma} \mathcal{L}_{ijrs}^{-1} \frac{\partial \phi}{\partial \sigma_{rs}^{(k)}} - \frac{(c^f)^2}{b^\sigma b^\gamma} \frac{\partial \sigma_y}{\partial \varepsilon^{p(k)}} \frac{\partial \phi}{\partial \sigma_{ij}^{(k)}} \\ \frac{(c^e)^2}{b^\sigma b^\gamma} \mathcal{L}_{klrs}^{-1} \frac{\partial \phi}{\partial \sigma_{rs}^{(k)}} - \frac{(c^f)^2}{b^\sigma b^\gamma} \frac{\partial \sigma_y}{\partial \varepsilon^{p(k)}} \frac{\partial \phi}{\partial \sigma_{kl}^{(k)}}, & \left(\frac{c^e}{b^\gamma} \right)^2 \frac{\partial \phi}{\partial \sigma_{rs}^{(k)}} \frac{\partial \phi}{\partial \sigma_{rs}^{(k)}} + \left(\frac{c^f}{b^\gamma} \right)^2 \frac{\partial \sigma_y}{\partial \varepsilon^{p(k)}} \frac{\partial \sigma_y}{\partial \varepsilon^{p(k)}} \end{array} \right]. \quad (27)$$

Given the scaled model problem, a method to find the desired scaled step vector is needed. The established dogleg method is utilized for this purpose and will be briefly reviewed here. For details and more extensive discussion, please see the text of Nocedal and Wright [37].

In order to iteratively update the solution vector, an initial, $\tilde{\Delta}^0$, and maximum, $\bar{\Delta}$, trust-region radius must first be defined. To this end, it is noted that the proposed algorithm is an incremental constitutive formulation. Therefore, it is expected that (i) maximum stress increment magnitude is equal to the magnitude of the difference between the trial and previously converged stresses and (ii) the maximum possible plastic strain increment would be the total strain increment. Therefore, the maximum trust-region radius, $\bar{\Delta}$, is set to,

$$\bar{\Delta} = b^\sigma \sqrt{(\sigma_i^{tr} - \sigma_i^n)(\sigma_i^{tr} - \sigma_i^n)} + b^\gamma d\bar{\varepsilon}, \quad (28)$$

where σ_i^{tr} and σ_i^n are the principal components of the corresponding stress states and

$$d\bar{\varepsilon} = \sqrt{\frac{2}{3} d\varepsilon_{ij}^{n+1} d\varepsilon_{ij}^{n-1}}. \quad (29)$$

The initial trust-region radius is selected as $\tilde{\Delta}^0 = \bar{\Delta}$. Importantly, this selection allows for a single step solution to be found if it exists.

To iteratively update the scaled step vector, $\{\tilde{\mathbf{p}}\}^{(k)}$, and scaled trust-region radius, $\tilde{\Delta}^{(k)}$, the aforementioned dogleg method is used. This approach is schematically represented in Fig. 2. Specifically, the dogleg method utilizes two scaled step vectors – the Cauchy point, $\{\tilde{\mathbf{p}}^c\}$, and full step, $\{\tilde{\mathbf{p}}^f\}$ – and finds the point running between them that lies closest to the edge of the trust-region. The former corresponds to the solution of the linear constrained (in terms of solution step magnitude) problem while the latter is the minimizer of the unconstrained quadratic model problem, Eqn. 23, and is essentially a scaled Newton-Raphson step. These vectors are given as,

$$\{\tilde{\mathbf{p}}^c\}^{(k)} = -\tau^{(k)} \left(\frac{\tilde{\Delta}^{(k)}}{\|\{\tilde{\mathbf{g}}\}^{(k)}\|} \right) \{\tilde{\mathbf{g}}\}^{(k)}, \quad (30)$$

with $\tau^{(k)}$ being,

$$\tau^{(k)} = \min \left[1, \frac{\|\{\tilde{\mathbf{g}}\}^{(k)}\|^3}{\tilde{\Delta}^{(k)} \{\tilde{\mathbf{g}}\}_{(k)}^T [\tilde{\mathbf{B}}]^{(k)} \{\tilde{\mathbf{g}}\}_{(k)}} \right], \quad (31)$$

and

$$\{\tilde{\mathbf{p}}^f\}^{(k)} = -[\tilde{\mathbf{B}}]_{(k)}^{(-1)} \{\tilde{\mathbf{g}}\}^{(k)}. \quad (32)$$

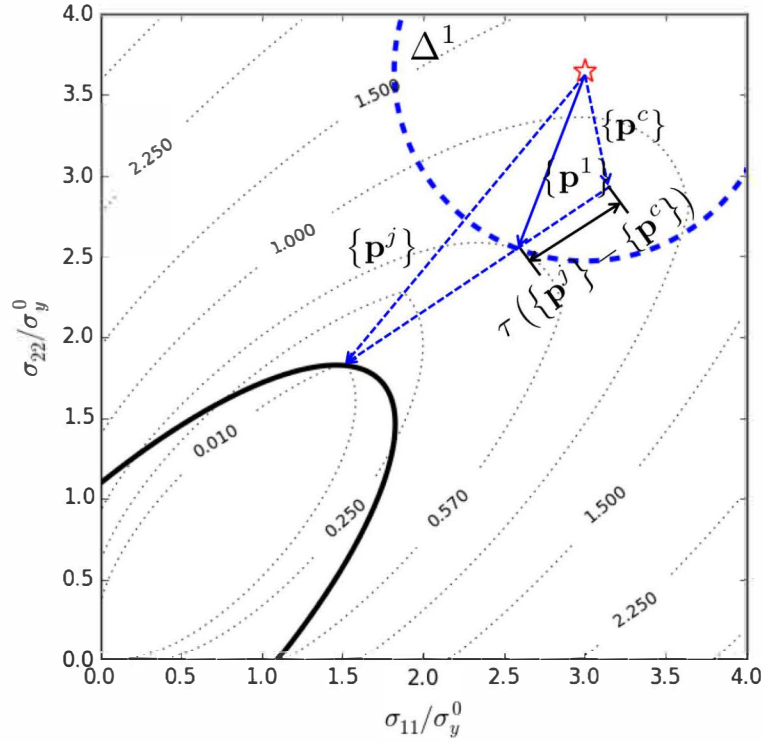


Figure 2. Illustration of the dogleg approximation to find $\{\mathbf{p}^1\}$ as a combination of the Cauchy point, $\{\mathbf{p}^c\}$ and full step vectors, $\{\mathbf{p}^j\}$.

If the Cauchy point, $\{\tilde{\mathbf{p}}^c\}^{(k)}$, lies on the boundary of the trust-region it is accepted as the minimizing step vector. When this condition is not satisfied, a vector running from the current material state (stress state in Fig. 2) to a point lying along the path connecting the Cauchy point and the full step with a magnitude of $\tilde{\Delta}^{(k)}$ is searched for. If the full step lies within the trust-region it is used. This process is given as,

$$\begin{aligned} &\text{if } \|\{\tilde{\mathbf{p}}^c\}^{(k)}\| = \tilde{\Delta}^{(k)} \\ &\quad \{\tilde{\mathbf{p}}\}^{(k)} = \{\tilde{\mathbf{p}}^c\}^{(k)} \\ &\text{else} \\ &\quad \{\tilde{\mathbf{p}}\}^{(k)} = \{\tilde{\mathbf{p}}^c\}^{(k)} + \tau \left(\{\tilde{\mathbf{p}}^j\}^{(k)} - \{\tilde{\mathbf{p}}^c\}^{(k)} \right), \end{aligned} \quad (33)$$

with $\tau \in (0, 1]$. To find τ , a simple bisection algorithm is employed in this effort in lieu of more complex approaches that have been adopted (*i.e.* conjugate gradient).

To update the trust-region radius, a measure of the iterative improvement is first determined. Specifically, $\rho^{(k)}$ is defined as the actual improvement over the expected improvement and is given as,

$$\rho^{(k)} = \frac{\psi(\{\mathbf{x}\}^{(k)}) - \psi(\{\mathbf{x}\}^{(k)} + \{\mathbf{p}\}^{(k)})}{\tilde{m}^{(k)}(\{\mathbf{0}\}) - \tilde{m}^{(k)}(\{\tilde{\mathbf{p}}\}^{(k)})}. \quad (34)$$

The updated scaled trust-region radius, $\tilde{\Delta}^{(k+1)}$, is determined based on this improvement measure. If substantial improvement is noted, the radius is increased. On the other hand, when insufficient gains are made the trusted region is reduced in size. The update process is chosen to be,

$$\begin{aligned}
& \text{if } \rho^{(k)} < \frac{1}{4} \\
& \quad \tilde{\Delta}^{(k+1)} = \frac{1}{4} ||\{\tilde{\mathbf{p}}\}^{(k)}||, \\
& \text{else} \\
& \quad \text{if } \rho^{(k)} > \frac{3}{4} \text{ and } ||\{\tilde{\mathbf{p}}\}^{(k)}|| = \tilde{\Delta}^{(k)} \\
& \quad \quad \tilde{\Delta}^{(k+1)} = 2\tilde{\Delta}^{(k)} \\
& \quad \text{else} \\
& \quad \quad \tilde{\Delta}^{(k+1)} = 2||\{\tilde{\mathbf{p}}\}^{(k)}|| \\
& \quad \quad \tilde{\Delta}^{(k+1)} = \min \left(\tilde{\Delta}^{(k+1)}, \tilde{\Delta} \right).
\end{aligned} \tag{35}$$

At this stage, an additional check is performed to ensure that the updated solution vector is actually an improvement. Specifically, if $\rho^{(k)}$ is above a threshold value, $\rho^{(k)} > \eta = 0.1$, the solution is considered to be an acceptable improvement. If not, the scaled step vector is unacceptable and rejected. The conditions for these checks are,

$$\begin{aligned}
& \text{if } \rho^{(k)} > \eta \\
& \quad \text{Accept Solution } \{\mathbf{x}\}^{(k+1)} = \{\mathbf{x}\}^{(k)} + \{\mathbf{p}\}^{(k)}, \\
& \text{else} \\
& \quad \text{Reject Solution } \{\mathbf{x}\}^{(k+1)} = \{\mathbf{x}\}^{(k)}, \\
& \quad \text{Force } (k+1) \text{ iteration to use the Cauchy step: } \{\tilde{\mathbf{p}}\}^{(k+1)} = \{\tilde{\mathbf{p}}^c\}^{(k+1)}.
\end{aligned} \tag{36}$$

Note, the updating procedures in Eqns. 35 and 36 largely follow those in Algorithm 11.5 of [37], but two key differences are evident in the proposed schemes versus that of [37]. The first being that in the event of a rejected step, the subsequent iteration is forced to take a Cauchy step. This is analogous to non-linear conjugate gradient algorithms taking a steepest descent step when unacceptable orthogonality is observed. Second, under acceptable conditions the trust-region radius scales with the step vector magnitude ($\tilde{\Delta}^{k+1} = 2||\{\tilde{\mathbf{p}}\}^k||$). These changes were found during algorithm development to improve convergence under numerically challenging conditions and as such are utilized here.

3. RESULTS

To investigate the performance of the the implementation presented in Section 2, it was implemented in the non-linear, quasistatics finite element code Sierra/SM [30]. First, a series of boundary value problems are considered in Section 3.1 to both verify the proposed algorithm and demonstrate its capabilities. The robustness and algorithmic performance are then extensively explored in Section 3.2. In both these cases, the proposed implementation will be compared to those of both a standard NR and a LS-NR algorithm. Finally, in Section 3.3, the impact of the scaling terms in Eqns. 13 and 22 will be explored. For these studies, properties representative of an elastic-plastic metal will be used ($E = 200$ GPa, $\nu = 0.3$, $\sigma_y^0 = 200$ MPa).

3.1. Verification Problems

A series of global boundary value problems are solved in this section using the three considered algorithms to verify the TR approach. For these verification problems, linear hardening (Eqn. 8) with $K = 20$ GPa will be considered along with a yield exponent of $a = 8$ that is common for FCC metals [50].

3.1.1. Biaxially Loaded Plate The capabilities of the TR numerical implementation developed in this work are now explored. For these investigations, a thin sheet ($1 \times 0.5 \times 0.01$ mm length (L) by width (W) by thickness (t)) comprised of nine linear hexahedral elements is considered. A set of six biaxial loadings corresponding to tension, compression, and various mixed conditions as shown in Fig. 3a are imposed along the indicated edges in Fig. 3b and analyzed. Eighth-symmetry, plane stress conditions are assumed for the remaining boundaries and fifty load steps are used for each analysis. The in-plane stress and strain results are presented in Fig. 4 for the NR, LS-NR, and TR solution methods.

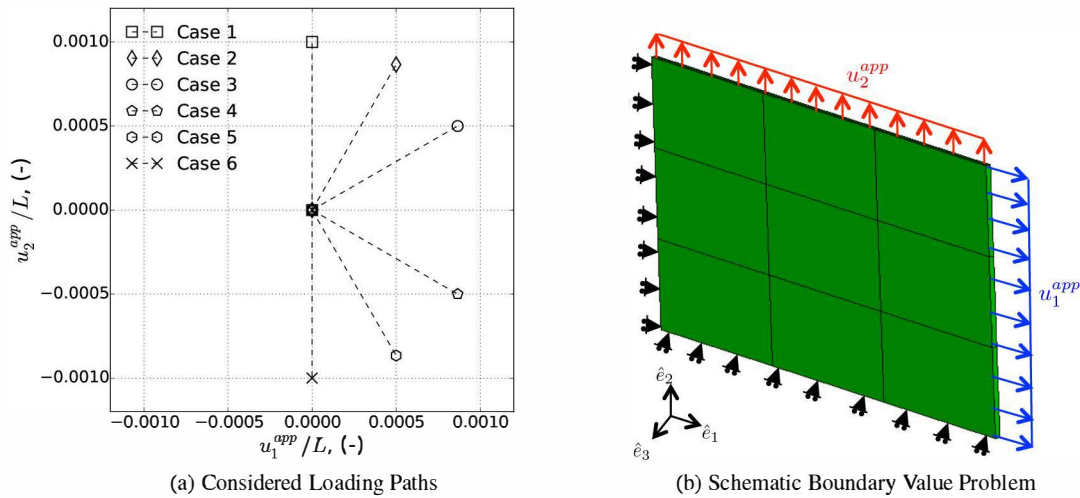
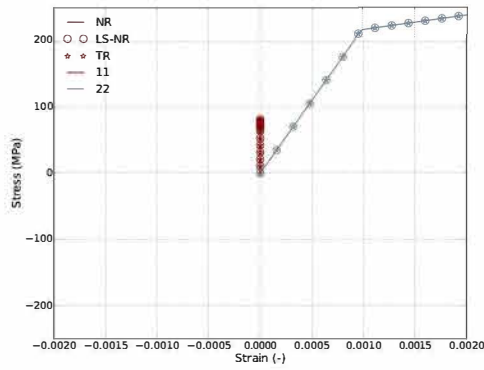


Figure 3. Considered (a) loading paths and (b) boundary value problem for the biaxially loaded plate tests

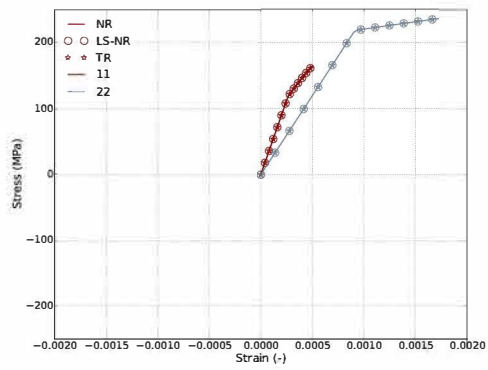
As is expected for a verification exercise, the three different implements produce the same results for the six responses of Fig. 4. The maximum relative differences of any of the algorithms was on the order of $1 \times 10^{-3}\%$ although the error was typically zero. In this way, the capability of the proposed trust-region model is demonstrated through a variety of simple loading paths. With respect to the speed of the different approaches, Table I presents the relative wallclock time of the different methods to those of the NR approach. This baseline is selected as it represents the most common implicit implementation and the relative cost of going to these more complex methodologies can be ascertained. Each case is run on a single processor of the same dedicated machine to minimize any variability and take just under a second. During plastic loading, two correction iterations are typically needed for each implementation. As can be seen from Table I utilization of the LS-NR or TR methods comes at only a modest cost. For the LS-NR approach, the additional time is less than 4% in all cases. The TR algorithm is typically slower than LS-NR approach but the overall increase in time is still less than 10% and in most cases approximately 5%.

Case	NR	LS-NR	TR
1	1.000	1.026	1.057
2	1.000	1.024	1.055
3	1.000	1.006	1.050
4	1.000	1.039	1.066
5	1.000	1.021	1.047
6	1.000	1.009	1.096

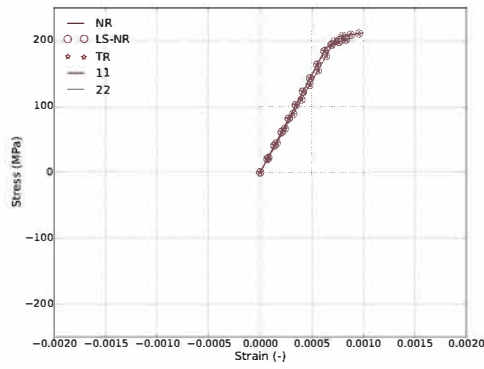
Table I. Relative (to the NR method) computational times for NR, LS-NR, and TR methods through the the six loading paths in Fig. 3a.



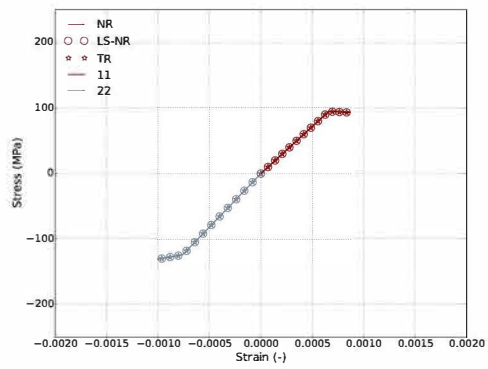
(a) Case 1 Loading Path



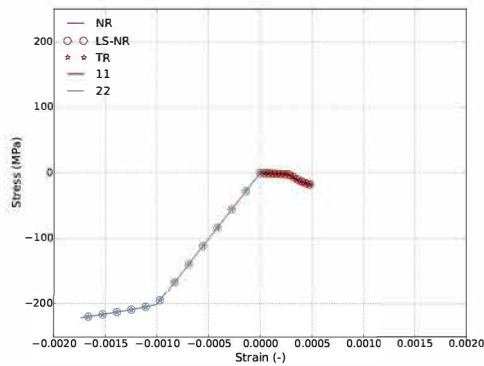
(b) Case 2 Loading Path



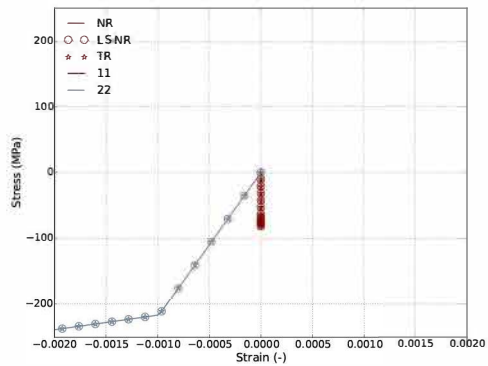
(c) Case 3 Loading Path



(d) Case 4 Loading Path



(e) Case 5 Loading Path



(f) Case 6 Loading Path

Figure 4. Stress-strain responses in the 11 and 22 directions for the six loading cases presented in Fig. 3a as determined by the NR, LS-NR, and TR methods.

3.1.2. Rod in Combined Tension and Shear The previous study focused on a relatively simple geometry of limited size. To further test the capabilities of this implementation, the problem of a rod subject to tension and shear previously tackled by Shterenlikht and Alexander [45] is considered. This specific problem is investigated for two reasons. First, as pointed out by Shterenlikht and Alexander, such a loading results in a variety of local loading paths testing a wide set of responses.

Second, in their study severe scaling issues prevented the simulations using the dogleg method from completing [45]^{||}. Although in that case the GTN constitutive model was used, consideration of this problem tests the ability of the extra scaling terms to address this issue and provides a challenge for the proposed implementation.

The rod in question is 100 mm long ($L = 100$ mm) with a circular cross section of radius 20 mm ($r = 20$ mm). All degrees of freedom along the top edge ($x_3 = L$) are fixed ($u_1 = u_2 = u_3 = 0$) while the bottom ($x_3 = 0$) is constrained to remain planar ($u_3 = 0$). To load the rod, a horizontal displacement of $u_1^{app} = 50$ mm is applied along the bottom, $x_3 = 0$, face. Schematically, these conditions are presented in Fig. 5a. The geometric origin is at the center of the bottom face. Two additional points (pt. $A = (-r/2, 0, 0)$ and pt. $B = (r/2, 0, 0)$) are indicated for subsequent analysis. For these simulations a fixed number of loading increments (200) is used and no global timestep cutbacks are allowed to better enable direct comparisons between the various results.

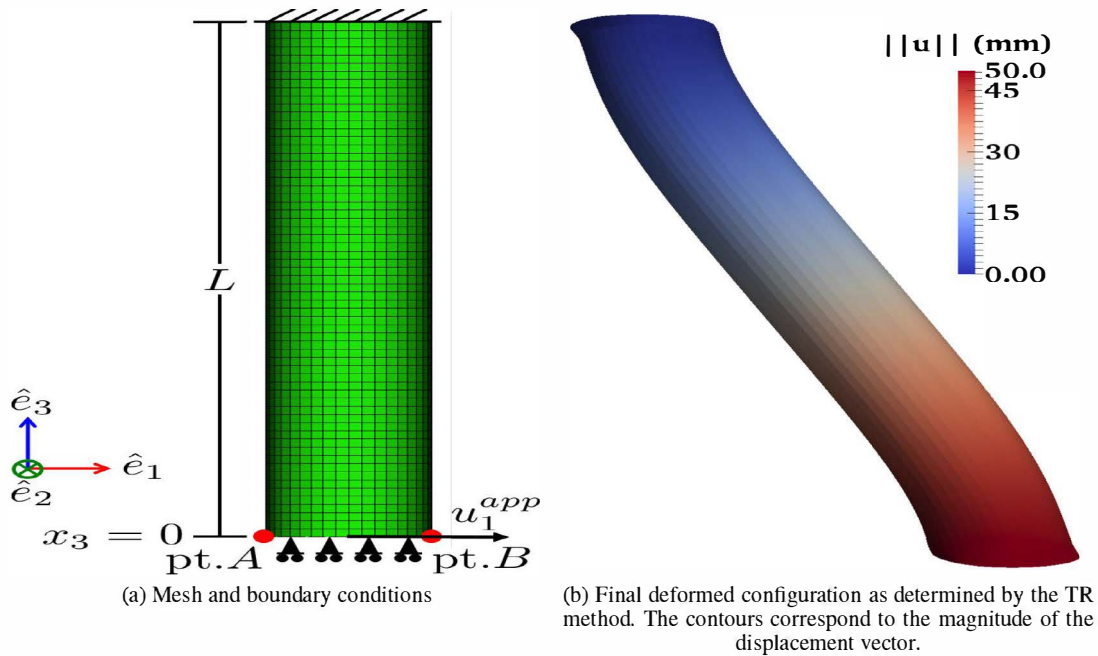


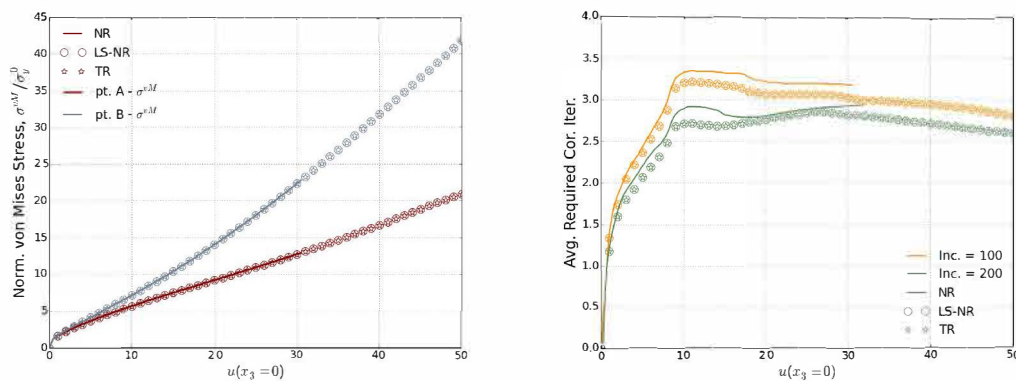
Figure 5. Summary of rod under combined tension and shear loadings: (a) Boundary conditions and mesh and (b) Final result.

Results of the analyses are given in Figs. 5b and 6 and good agreement is noted between the various numerical implementations *that converge*. Specifically, the TR and LS-NR results are sufficiently robust such that local problem may be solved even under large, complex loadings enabling the global solution to be found. The same cannot be said for the NR method as later loadings prove too challenging for the scheme and the problem fails to converge. This capability of the TR scheme is further demonstrated in Fig. 5b which shows the final configuration at the end of the loading highlighting the substantial deformations in this case. Additionally, Fig. 6a presents the effective von Mises stress at points A and B , respectively, at different applied displacements. No measurable difference is found at either point between the LS-NR and TR methods at any applied loading further verifying the proposed implementations. Agreement is also observed with the NR method through the portions of the simulation that are successful.

^{||} In their study [45], the Levenberg-Marquardt approach was able to successfully analyze this problem. However, as the dogleg algorithm they discuss more closely resembles the current method, the performance of that approach is of more concern for motivation and comparison purposes.

To consider the relative performance and speed of the algorithms, the average number of correction iterations needed at each load step, averaged over the entire model, is determined and plotted in Fig. 6b. The average number of iterations is plotted in lieu of the maximum to better assess the overall performance of the algorithms rather than extremum cases. As can be seen in the results of Fig. 6b identical responses are noted for the LS-NR and TR implementations. Both of these approaches require fewer correction iterations than the more standard NR – highlighting additional benefits for these methodologies in terms of reducing computational cost.

As fixed time incrementation is used in these simulations, it is possible that the selected timestep sizes may influence performance of the various algorithms. To investigate this possibility, a second set of simulations utilizing only 100 loading increments are performed and the results presented in Fig. 6b. As expected, a slightly higher average number of correction iterations is required versus the 200 increment case. The same trends in comparing the three methods are also observed in this case highlighting the improved performance of the LS-NR and TR methods over the more traditional NR case. With respect to speed, no appropriate comparison is available between the NR and other methods due to lack of convergence in the former. In comparing the LS-NR and TR results, it is noted that the TR cases had 3.3% and 3.8% higher wallclock times than the LS-NR approach for the 100 and 200 increment analyses, respectively. Although slightly more expensive, these timings again show comparable speed performance between the two implementations.



(a) Effective von Mises stress at pt. A and B, respectively. (b) Element averaged inelastic correction iterations required utilizing different load incrementation.

Figure 6. Results of rod tension/shear problem: (a) von Mises effective stress and (b) required correction iterations over loading as determined by the NR, LS-NR, and TR methods.

3.2. Numerical Robustness

With the capabilities of the trust-region implementation verified through different boundary value problems, the next issue to consider is that of the robustness of the proposed algorithm. To that end, the methodology of Scherzinger [29] is adopted. Through this approach, a series of trial stress states are determined and used as input to the considered algorithm. The number of iterations needed to converge (or lack thereof) is recorded. The considered initial stress states scan the π -plane between the initial yield surface and a surface whose equivalent effective stress is 30 times that of yield ($\phi = 30\sigma_y^0$). Although quite large, $30\sigma_y^0$ is selected as strain increments may be produced during inelastic deformation resulting in quite substantial trial stresses. Given that local convergence failure can lead to global time step cutbacks and potentially noticeable time costs, robustness under large trial stresses is desired. Additionally, in FE implementations using a preconditioned conjugate gradient solver (like Sierra/SM [30]) substantial variation in the displacement (strain) field during global iteration may be observed further necessitating robustness at seemingly excessive trial stress

states. Sampling in this fashion results in over 95,000 trial stress states whose numerical response is considered.

First, three different yield exponents are considered ($n = 6, 8, \text{ and } 100$) that sequentially increase the curvature of the yield surface. The former values are commonly used for BCC and FCC metals, respectively [50], while the latter $n = 100$ case approximates the Tresca yield surface and its numerically problematic corners. Trivial cases of $n = 2$ and 4 corresponding to a von Mises yield surface are neglected as the results show a uniform, single step return mapping procedure as is expected for a radial-return. For these studies, perfect plasticity ($K = 0$) is considered as a previous study [29] indicated hardening had little impact on algorithmic performance. For the TR algorithm, one correction must be made for usage in this analysis. Specifically, as no strain increment or previous material state is given a modification is needed for $\bar{\Delta}$ in Eqn. 28. Therefore, following the motivations discussed regarding Eqn. 28, it is noted (i) the trial stress, not total strain increment, is used as input to the constitutive integration routine (ii) with $K = 0$ the effective stress measure of the solution stress will be equal to the yield stress and (iii) the total strain increment corresponds to the elastic strain of the trial stress state. Therefore, for this study, the modified maximum trust-region radius is written as,

$$\bar{\Delta} = b^\sigma \sqrt{(\phi(\sigma_{ij}^{tr}) - \sigma_y^0)(\phi(\sigma_{ij}^{tr}) - \sigma_y^0)} + b^\gamma \sqrt{\frac{2}{3} \mathbb{C}_{ijkl}^{-1} \sigma_{kl}^{tr} \mathbb{C}_{ijmn}^{-1} \sigma_{mn}^{tr}}. \quad (37)$$

The convergence maps for the case of yield exponents $n = 6, 8, \text{ and } 100$ are presented in Fig. 7. These maps present the number of correction iterations needed to achieve convergence at a given trial stress space with the results projected onto the deviatoric π -plane. Lighter colors denote fewer correction iterations and points marked in red do not converge in less than 40 iterations.

Importantly, between the three convergence maps in Fig. 7 it is observed that the TR algorithm converges for almost every case in less than 40 iteration. In fact, all considered trial stresses for the $n = 6$ and 8 cases do converge by the specified iteration. Some ($< 5\%$) points may be found in the $n = 100$ Tresca-like case that do not converge in less than 40 iterations. Furthermore, as expected the three maps exhibit a six-fold symmetry that is anticipated for the isotropic yield surface. Additionally, it can clearly be seen that as the exponent and corresponding curvature increases so do the number of required iterations. The $n = 6$ cases shows a fairly uniform light coloration indicating relatively easy convergence for most cases – even at substantial trial stresses. At the higher $n = 8$ value, some more computational difficulty arises in domains closer to the higher curvature areas. In the limit case of $n = 100$ a generally more expensive convergence may be observed. A more diffuse map is noted but a clear trend of increased correction iterations nearest the corners is evident. Additionally, the cases that do take more than 40 iterations to converge seem to occur at the edges of these domains.

To more extensively compare the results presented in Fig. 7, cumulative distributions of the total percentage of sampled cases that have successfully converged by a given correction iteration are presented in Fig. 8. Although the convergence maps for the NR and LS-NR cases are not presented in this work (see [29]), their responses were calculated and the corresponding cumulative convergence distributions are also presented in Fig. 8. It should be noted that given the sampling used to generate Fig. 7, the current distributions do not correspond to that expected during a typical boundary value problem.

From the results of Fig. 8, a number of important observations regarding the convergence characteristics of the three methods may be discerned. First, in all three cases, the response of the different methods through the first four correction iterations is quite similar. At this point, the converged cases of the LS-NR approach increase at a quicker rate than the other two cases. Eventually, for all three exponents, the TR implementation starts to catch up to the LS-NR approach and in the $n = 8$ case actually achieves a higher percentage of converged trial stresses. At this stage, the number of cases converging per iteration decreases and both the LS-NR and TR method approach complete convergence (100% cumulative convergence). The NR implementation, however, cannot achieve complete convergence and fewer cases converge with larger exponents. Importantly, for all of the considered yield surfaces the LS-NR and TR method always converge. In

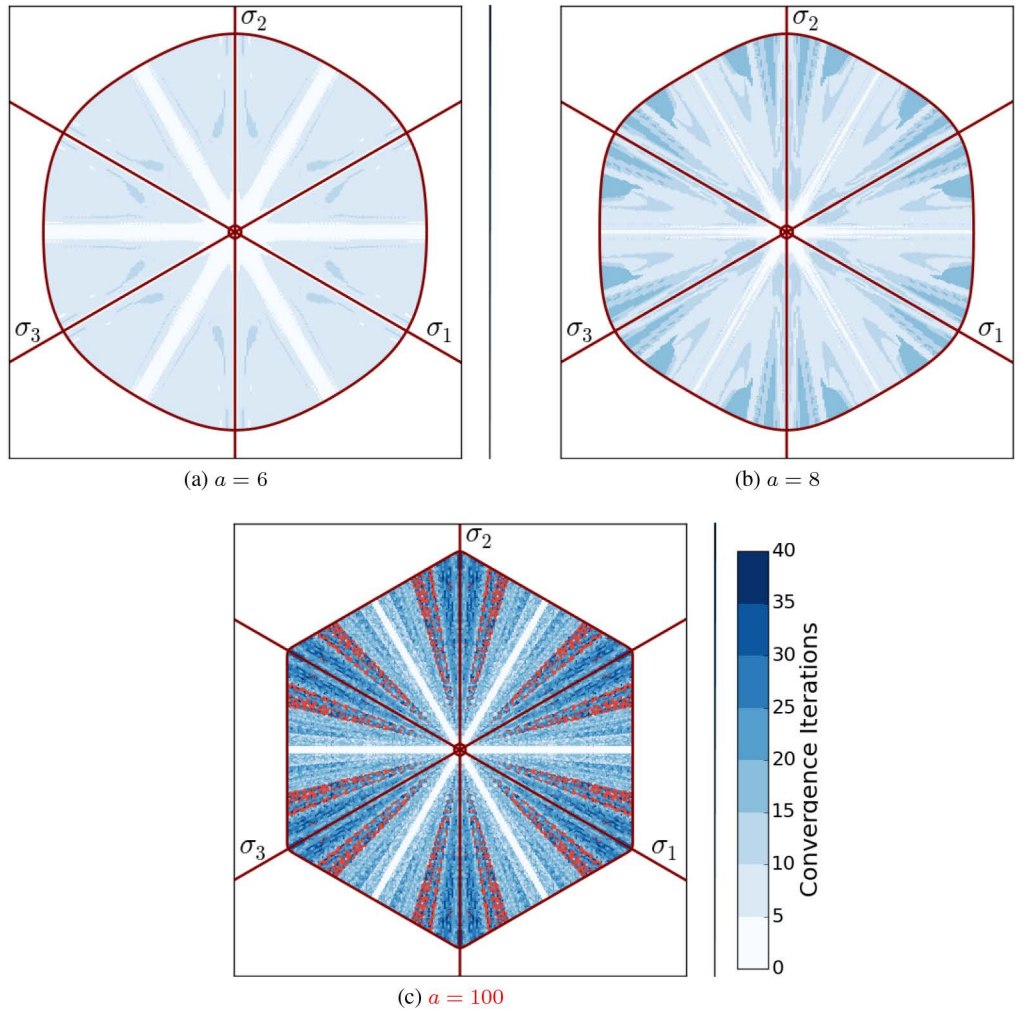


Figure 7. Convergence maps for the TR algorithm giving the number of correction iterations needed at a trial stress state in deviatoric space to achieve convergence. Three Hosford yield surface with different exponents, a , are considered. The same scale is used for all figures and points that do not converge within 40 iterations are marked red.

the $a = 100$ case this does not always occur by the 40th iteration, but it does occur within 65. These problematic trial stresses represent **less than 5%** of the tested cases. To summarize the capabilities of the different numerical schemes, the number of iterations needed to achieve complete, 100% cumulative convergence are given in Table II. As would be expected, increase in the curvature of the yield surface (reflected in higher values of a) leads to more iterations required to achieve complete convergence.

a	NR	LS-NR	TR
6	-	11	13
8	-	24	20
100	-	45	64

Table II. Number of iterations required to achieve complete, 100% cumulative convergence for different Hosford yield exponents (a) for the NR, LS-NR, and TR approaches.

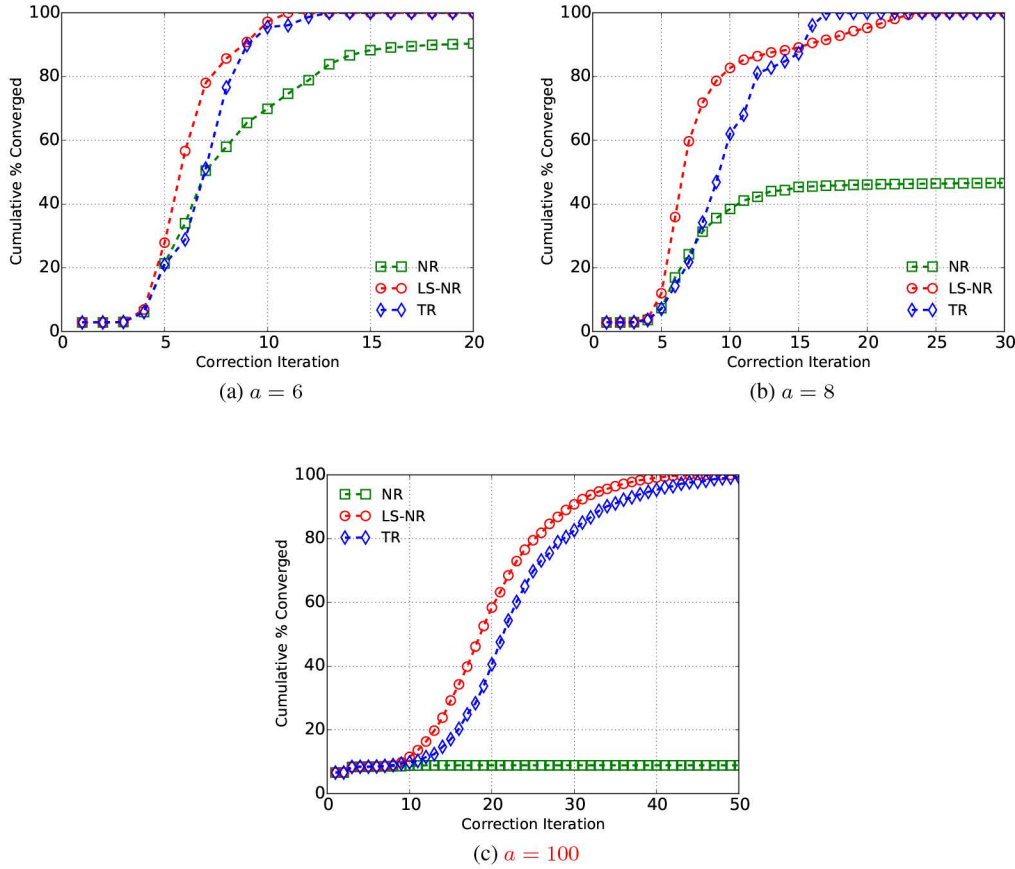


Figure 8. Cumulative convergence distributions of NR, LS-NR, and TR methods for three different Hosford yield surface exponents, a . Considered cases come from sampling the π -plane at locations whose scalar effective stresses are $\sigma_y < \phi(\sigma_{ij}^{tr}) \leq 30\sigma_y$.

Based on these results, the TR performance is clearly superior to the NR method in terms of robustness. With respect to the LS-NR method, such a distinction is less clear. Figure 9 explores such a comparison by presenting the iteration difference between the two approaches. Specifically, in Fig. 9a a π -plane map in which the iteration change in going to the TR approach from that of the LS-NR is plotted for the $a = 8$ case. Put another way, the iteration change is equal to the iterations needed for the TR method minus those of the LS-NR approach. As such, trial stress states in red indicate regions in which the TR algorithm requires more correction iterations than the LS-NR and blue indicates areas in which the TR approach requires fewer correction iterations and therefore exhibits superior performance. Regions in green denote cases in which the TR required at least ten fewer correction iterations while purple is used for trial stress states needing at least ten more. In terms of extremes, the proposed TR scheme required eighteen fewer or fourteen additional correction iterations versus the LS-NR. From the results of Fig. 9, it may be observed that the two algorithms show nearly identical convergence behaviors in domains with smaller effective stress measures. At trial stresses with larger effective stresses, substantial gains (in terms of needed iterations) may be obtained by using the TR method in some shear dominated domains. Figure 9b expands on this by presenting the average change in correction iterations for a given effective stress measure for all three yield exponents. Averaging in this fashion samples a variety of stress states (e.g. shear-dominated; uniaxial) and yields a representative metric for a given stress level. From such an analysis, it is observed that at lower stress measures there is no difference in the required number of

iterations. As the yield exponent n increases, however, the length of this similarity decreases. For the smallest exponent considered, the LS-NR algorithm outperforms (in terms of correction iterations) the TR at all effective stress levels. At the larger $n = 8$ value, regions in which both TR and LS-NR algorithm exhibit superior performances are clearly observed. When the curvature approaches that of Tresca ($n = 100$), large oscillations are seen in the iteration differences. Nonetheless, the trend of the response is that **LS-NR** seems to perform better than the **TR** at most effective stress levels. **As the average differences between the integration schemes is reasonable**, these results indicate the TR algorithm may hold promise for more computationally challenging problems.

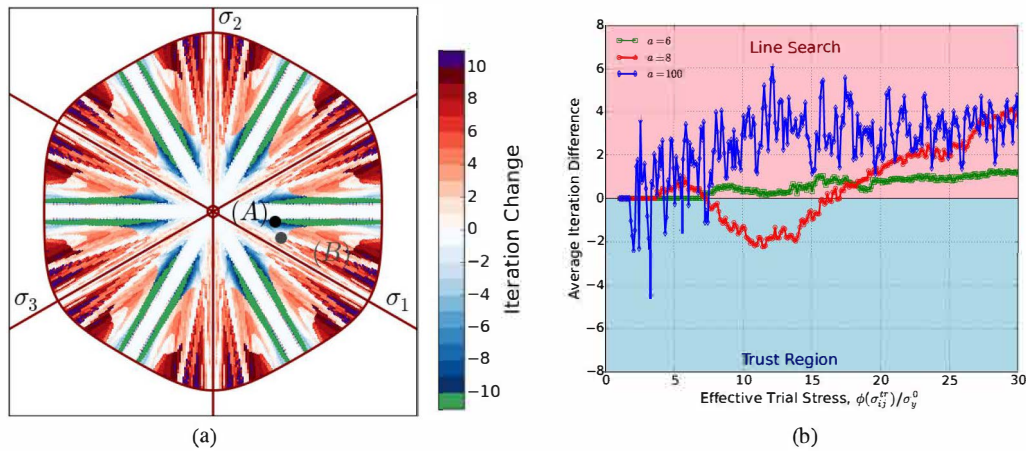


Figure 9. Iteration change ($\text{iter}(\text{TR}) - \text{iter}(\text{LS-NR})$) needed by the TR algorithm versus the LS-NR approach: (a) π -plane map in which any trial stress state requiring 10 or more fewer iterations is colored green while any case needing 10 or more additional iterations is shaded purple; (b) average number of correction iterations needed for a given effective stress measure. The results correspond to the $n = 8$ Hosford yield surface and two specific stress states in (a) are labelled (A) and (B) for later analysis.

To consider the convergence characteristics and response of the TR algorithm more closely, two trial stresses are marked in Fig. 9a. The first, marked (A), corresponds to a trial stress state, $\sigma_{ij}^{tr(A)}$, with principal components $\sigma_1 = 1,378$ MPa, $\sigma_2 = -242$ MPa, and $\sigma_3 = -1,135$ MPa and is one of lower effective trial stress states showing a more than 10 iteration improvement in going to the TR method. The second, marked (B), is for a trial stress state, $\sigma_{ij}^{tr(B)}$ whose principal components are $\sigma_1 = 2,187$ MPa, $\sigma_2 = -876$ MPa, and $\sigma_3 = -1,311$ MPa and is selected for the opposite reason. In this case, the TR implementation required **seven** more iterations. Figures 10 and 12 explore the source of these differences by presenting the return mapping (convergence) trajectories. These trajectories start from the trial stress state (denoted as the “0” iteration) and plot the successive stresses that are found during the return mapping process. In this way, the path taken during the inelastic correction process may be visualized to compare and analyze the performance of the different algorithms.

In Fig. 10, the return mapping processes of the NR (10a), LS-NR (10b), and TR (10c) algorithms with a trial stress of $\sigma_{ij}^{tr(A)}$ are presented. Figure 11 gives the evolution of the merit function for the LS-NR and TR approaches along with the scaled trust-region radius during inelastic correction. From the results of these two figures, it is observed that the return path of the TR implementation differs from the other two in the very first iteration. Specifically, the NR and LS-NR both take a substantial step that brings the stress state much closer to the yield surface but overshoots in terms of flow direction. This leads to a second step that attempts to correct the direction but again goes too far. The LS-NR algorithm is eventually able to find the correct direction and descend to the yield surface using a large number of step size cutbacks (see [29]). In Fig. 11, this behavior appears as the set of iterations with only small changes to the merit function. As the NR approach cannot cutback, it is unable to compensate and cannot converge. The TR approach, on the other-hand, actually rejects

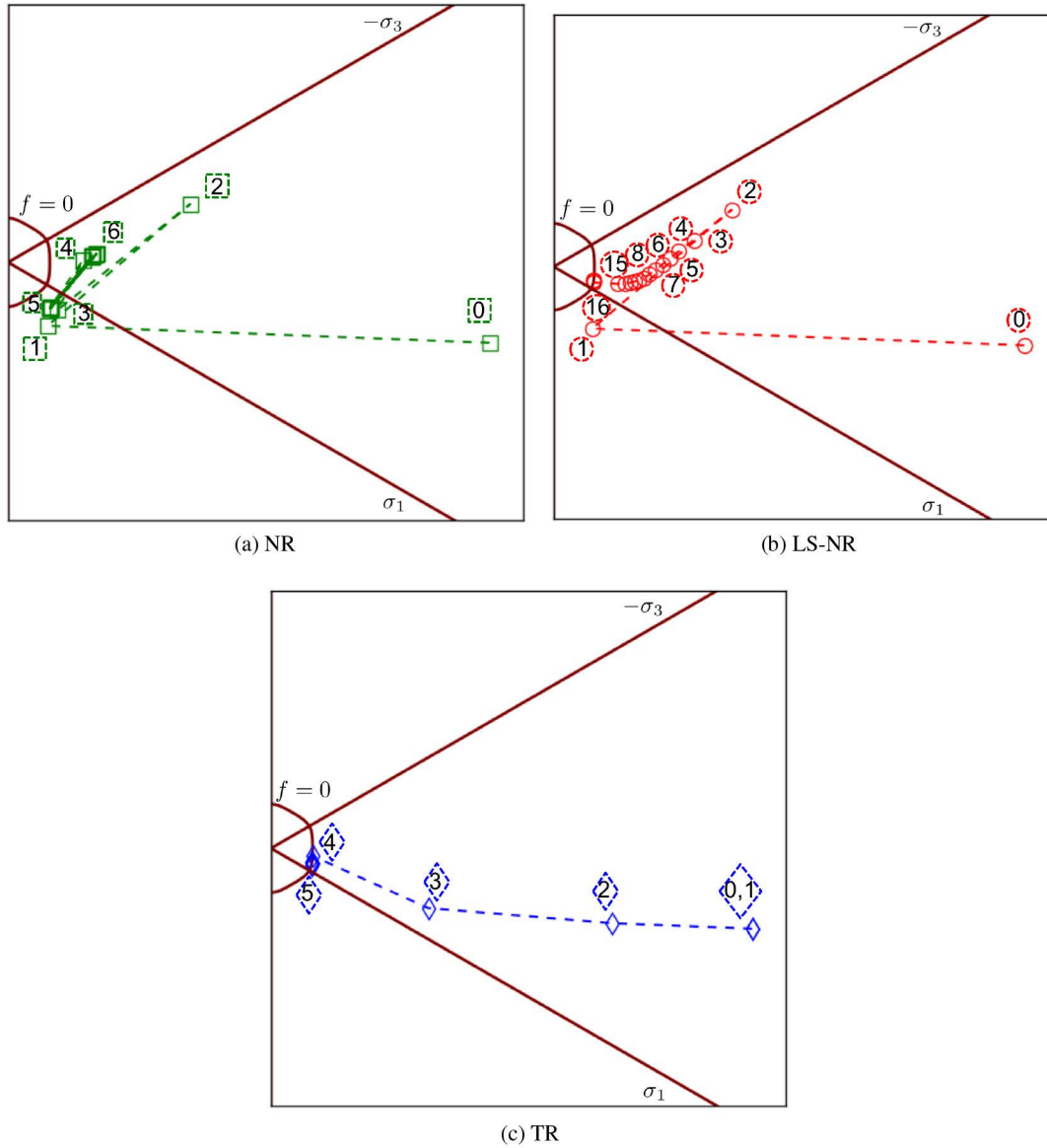


Figure 10. Inelastic correction process of the CPP-RMA problem solved by the (a) NR, (b) LS-NR, and (c) TR algorithms with an initial trial stress of $\sigma_{ij}^{tr(A)}$. The marked points are used to indicate the current correction iteration for select points. As the NR algorithm does not converge, only the first 20 iterations are plotted.

this initial step and cuts back on the scaled TR radius as shown in Fig. 11. This leads to a much smaller second correction iteration for the TR that does not overshoot the desired flow direction. Subsequent steps then eventually find the correct direction and converge to the solution in far fewer iterations than the LS-NR approach.

Figures 12 and 13 present the return mapping processes of the the algorithms and merit function and scaled trust-region evolution, respectively, for the opposite case ($\sigma_{ij}^{tr(B)}$) in which the LS-NR (and NR) approaches outperform the TR. Initially, the return paths of the three methods are similar to that discussed for Fig. 10. The LS-NR and NR both take large initial steps while the TR implementation rejects the first, large step. Unlike the previous results, the second correction for the LS-NR and NR algorithms finds the correct direction and quickly descends to the solution.

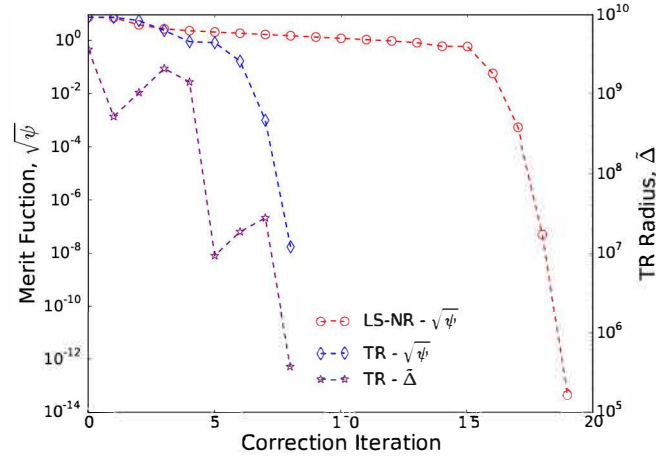


Figure 11. Merit function, ψ , and TR radius, $\tilde{\Delta}$ evolution through the LS-NR and TR inelastic correction process associated with an initial trial stress of $\sigma_{ij}^{tr(A)}$.

Similar to the previous case, the TR approach seeks to find the correct direction early and descend to the correct solution. As indicated in Figs. 12c and 13, however, this requires multiple iterations being rejected and TR radius cutbacks. This leads to more correction steps than the LS-NR and NR approaches. Interestingly, the fact that in both these cases the TR seeks to stay so close to the correct descent direction is likely indicative of the scaling introduced into the merit function.

3.3. Impact of Scaling

Trust-region methods and the performance of corresponding algorithms are known to be sensitive to scaling due to the large differences in magnitude of the solution variables. The size of this disparity is tied to the selection of units used in the analysis and as such careful consideration and selection of these characteristics may be able to minimize the impact and remove the necessity of the scaling constants. In many cases, however, other factors in the analysis (*e.g.* mesh size) may restrict the freedom in selecting the units and inhibit the ability of the analyst to address this issue without the utilization of normalization terms. As such, in this section the impact of these terms on algorithmic performance is investigated and assessed. To enable such considerations, two transformations were introduced – $[\mathbf{D}^1]$ in Eqn. 13 and $[\mathbf{D}^2]$ in Eqn. 22 – to address scaling in the merit function and state variable increments, respectively. Up to this point, these values have been fixed with limited motivation as to their selection. Therefore, to consider the impact of the relative contrast of merit function components, a modified weighting constant for the flow rule residual of the form $c^e = \beta (E / \sigma_y^0)$ is introduced with β being a constant. At this point, it is emphasized that β is neither a material model nor algorithmic parameter. Instead, it is an artificial scaling variable introduced to study the effect of the *relative* scaling of the two residual contributions. Furthermore, it is noted that as c^e and c^f are scaling quantities they themselves have different units. In fact, by comparing the expressions for c^e and c^f it is noted the two terms differ by E . As such, a base unit must be selected to examine these effects. For this study, the Pascal (Pa) is selected as the basis as casting quantities in this form provides the maximum contrast in residual contributions thereby producing a “worst-case” in terms of scaling that can best illustrate this subject. Therefore, noting that $E = 2.0 \times 10^{11}$ Pa, the range $10^{-11} \leq \beta \leq 10^0$ will be explored. As such, the larger values of β correspond more closely to the properly scaled cases studied throughout earlier sections of this work. Smaller artificial scalings lead to more substantial deviations from what has been used and essentially serve to decrease the contribution of the flow rule residual, r_{ij}^e , in the merit function

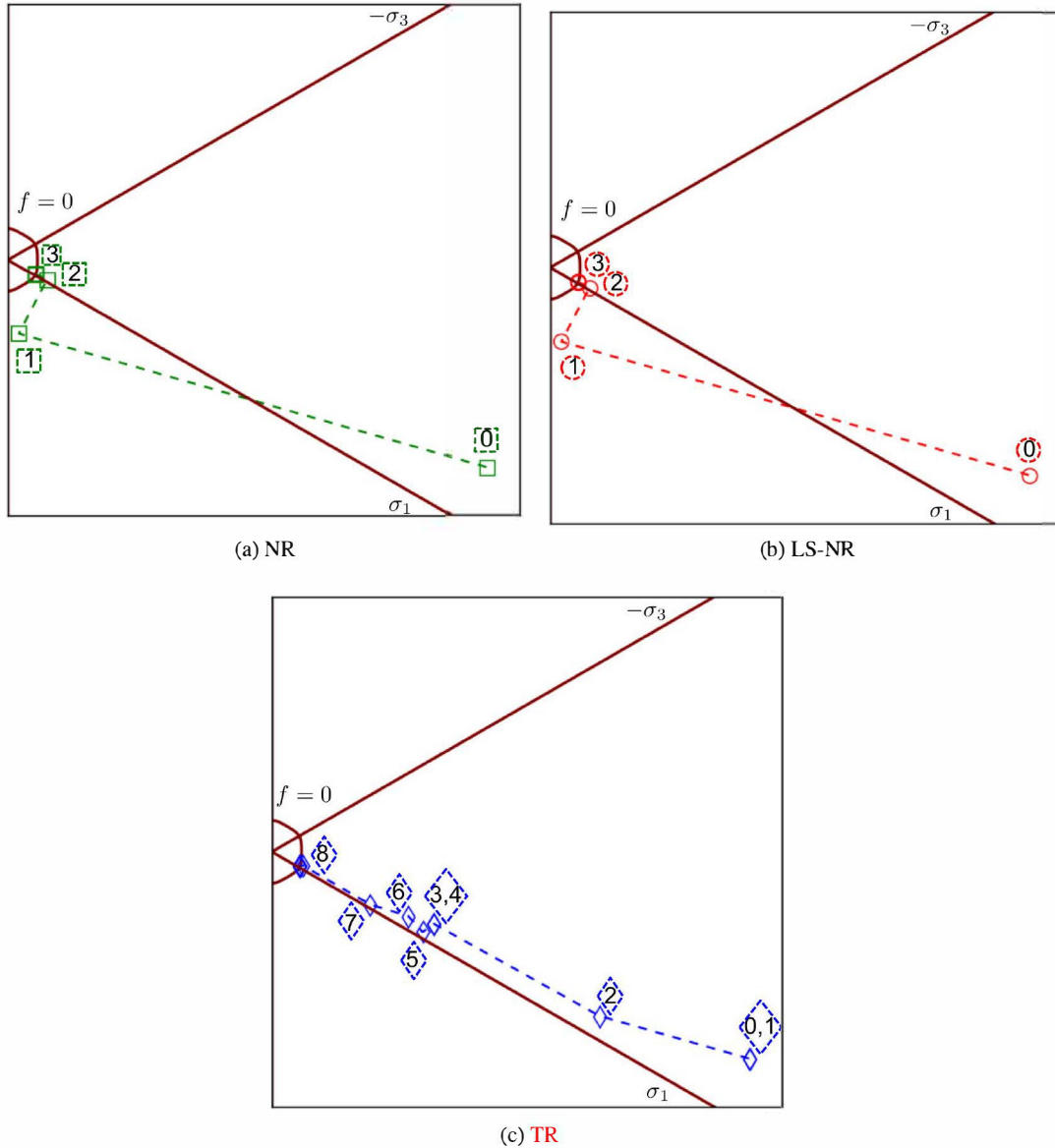


Figure 12. Inelastic correction process of the CPP-RMA problem solved by the (a) NR, (b) LS-NR, and (c) TR algorithms with an initial trial stress of $\sigma_{ij}^{tr(B)}$ (MPa). The marked points indicate the current correction iteration although for clarity not all points are labelled.

evaluation. In essence, $\beta = 10^{-11}$ approximates an unscaled merit function like that commonly used in many conventional implementations.

Figure 14 presents a summary of cumulative convergence distributions (like those in Fig. 8) determined for 100 different values of β . Specifically, for each artificial scaling the necessary number of iterations needed to first achieve *at least* cumulative convergences of 10%, 25%, 50%, 75%, and 100% are presented. If the specified threshold is not reached for a given value of β , it is not plotted. The percentage of unconverged states (after 100 iterations) is also presented to highlight any lack of convergence. As iterations occur only in discrete values, the actual cumulative convergence at these points will not match the given limit value exactly. From the results in Fig. 14, it can be observed that scaling and the relative contrast between the residual terms in the merit function play a strong role in the performance of the TR algorithm. For instance, only properly scaled

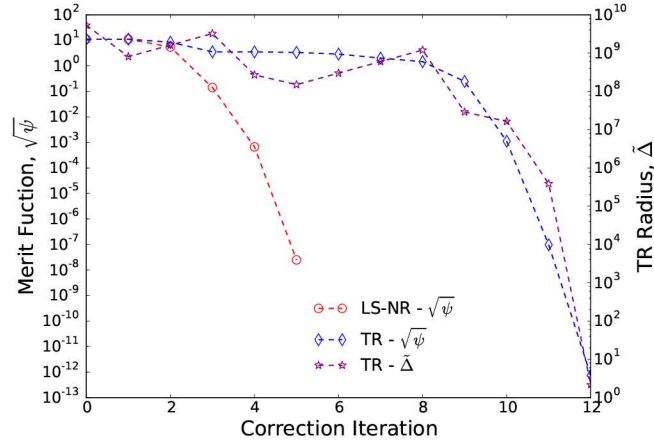


Figure 13. Merit function, ψ , and TR radius, $\tilde{\Delta}$ through the LS-NR and TR inelastic correction process associated with an initial trial stress of $\sigma_{ij}^{tr(B)}$.

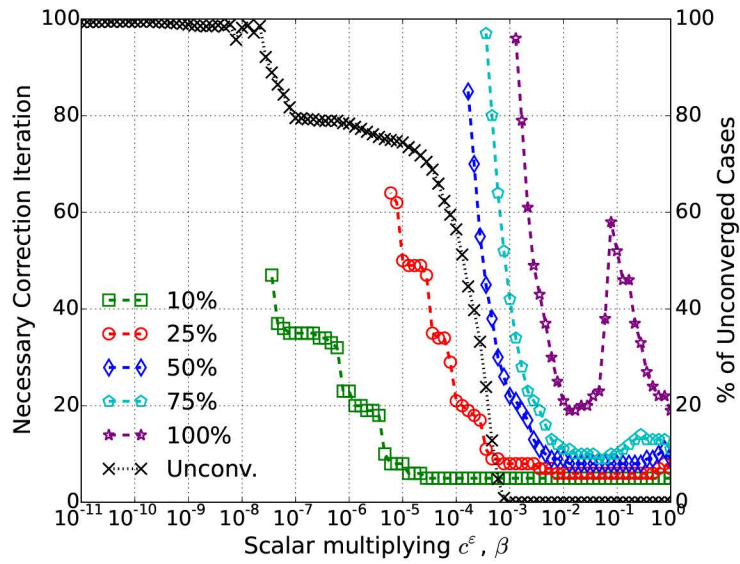


Figure 14. Influence of relative contrast between c^e and c^f on the convergence characteristics of the proposed TR method. Results correspond to the necessary number of correction iterations to achieve the specified cumulative convergence threshold as a function of the artificial scaling parameter, β , in which $c^e = \beta (E/\sigma_y^0)$.

cases ($\beta \gg 10^{-3}$) exhibit complete convergence while artificially scaled problems ($\beta \leq 10^{-4}$) converge for less than $\approx 40\%$ of the trial stress states and may be considered poorly scaled. As such, it can clearly be seen that careful selection of normalization constants is necessary for the TR implementation.

To investigate the source of the strong scaling dependence of the proposed approach, Fig. 15 presents the return mapping process of a case with no artificial scaling, $\beta = 10^0$, and one with moderate artificial scaling ($\beta = 10^{-4}$). The trial stress state in this study corresponds to $\sigma_{ij}^{tr(A)}$ presented in Fig. 10. For the artificially scaled response, the first 100 iterations are presented while

the $\beta = 10^0$ response corresponds to the TR results previously presented and discussed in Fig. 10c. In Fig. 15a, it can be observed that the the inappropriately scaled case ($\beta = 10^{-4}$) initially takes the large step bypassed by the correctly scaled case. This indicates the impact of the scaling on the algorithm performance as this large step is now acceptable in terms of improvement in the merit function. Subsequent iterations head straight to the yield surface albeit well away from the appropriate location. Given the poor scaling of the problem, the successive correction steps are unable to move away and instead oscillate around the surface **slowly** heading towards the correct solution (Fig. 15b). In this context, the decreased β leads to the consistency condition contribution dominating the model problem underlying the TR approach. Therefore, unless the merit function is properly scaled, the TR method cannot simultaneously satisfy the two conditions at the heart of the inelastic correction problem.

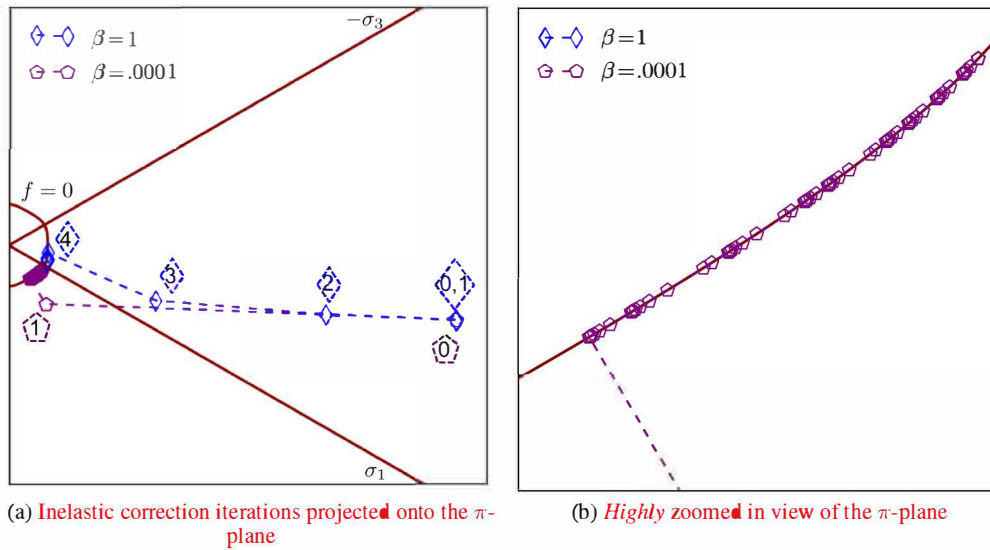


Figure 15. Inelastic correction process of trust-region (TR) method using two different stress normalization scalings for the $\sigma = 8$ yield surface with an initial trial stress of $\sigma_{ij}^{tr} = \sigma_{ij}^{tr(A)}$ (MPa). The results are presented in terms of the artificial scaling parameters, β , where $c^e = \beta (E/\sigma_y)$.

The second potential source of scaling is associated with the solution variables and is addressed via selection of b^σ and b^γ . In all previous cases, $b^\sigma = 1$ and $b^\gamma = 2\mu$ which, following the discussion and assumptions at start of this section, means a difference of roughly eleven orders of magnitude. To investigate this effect, the cumulative convergence thresholds utilizing a plastic consistency scaling of $b^\gamma = \beta 2\mu$ (with $10^{-11} \leq \beta \leq 10^0$) are determined as a function of the artificial scaling parameter, β , and the corresponding results are plotted in Fig. 16. In Fig. 16a, b^γ is scaled independently of the merit function to isolate the impact of the state variable terms.

From the results of Fig. 16a, a moderate dependence on the scaling of b^γ is observed. Specifically, while the consistency multiplier is still roughly stress measured ($b^\gamma \gtrsim 0.2\mu$), algorithmic performance remains relatively unchanged. Below this level, however, a substantial and rapid degradation is noted with the number of convergence iterations needed to achieve 75% and higher thresholds doubling over two decades. However, these responses eventually stabilize when $\beta \leq 10^{-5}$ and the necessary correction iterations to achieve the remaining convergence thresholds do not change. At the lower convergence thresholds (10%, 25%) scaling has very little impact as even in the worst case few extra iterations are needed. Additionally, convergence is achieved for more than 99% of the considered trial stress states regardless of the artificial scaling coefficient. This response is in contrast to the impact of the merit function scaling (Fig. 14) in which a pronounced effect is observed with β and very few cases are able to converge. Therefore, it is reasonable to

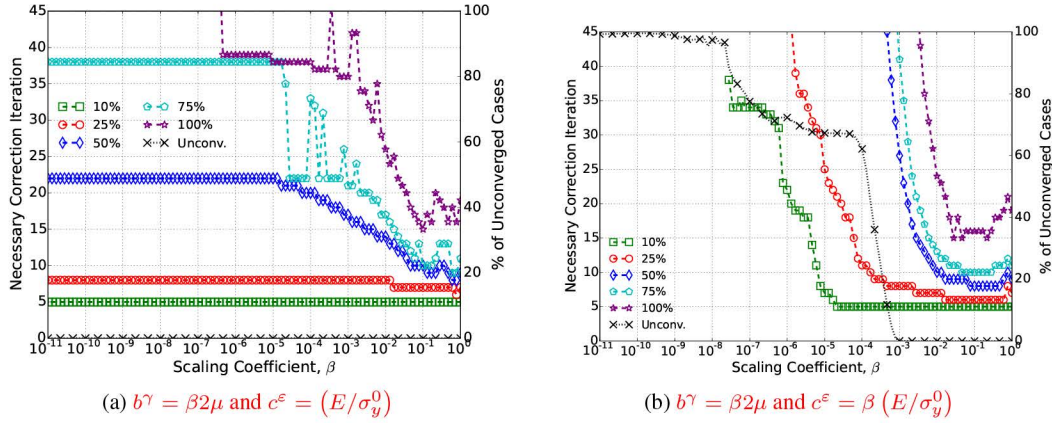


Figure 16. Influence of scaling the state variables on the convergence characteristics of the trust-region method in terms of the artificial scaling parameter, β .

state that although appropriate scaling of the state variable coefficients is necessary to achieved the desired **performance**, the dependence is far less severe than on the merit function constants.

Given the disparity between the scaling dependence of the merit function and state variables, one possibility to try and alleviate the convergence difficulties of the former is to scale both terms together. This course is investigated in Fig. 16b and the corresponding results are observed to much more closely follow the merit function scaling trends than those of b^γ . Specifically, complete convergence is only achieved when properly scaled ($\beta \gtrsim 10^{-3}$) and limited convergence is seen over the rest of the domain. Although similar to the results of Fig. 14 some differences may be observed. Specifically, the 25% convergence threshold is achieved over a larger β range and **the lack of an iteration increase around $\beta = 10^{-1}$ in the 100% case**. With respect to this latter point, no clear trend is observed. In any case, these results serve as further evidence that inappropriate scaling in the merit function leads to drastically reduced performance of the TR method. Additionally, scaling the state variables in a similar fashion cannot rectify this issue.

4. CONCLUDING REMARKS

In this work, a novel method utilizing the TR method for the implicit integration of plasticity models is presented. Unlike previous approaches, the non-linear equations forming the core of stress updating procedure are not solved via the Newton-Raphson method thereby bypassing the corresponding robustness issues and avoiding the use of substepping or line search techniques. The proposed methodology was discussed in detail and implemented for a Hosford plasticity model. Through a series of boundary value problems, the model response was verified against traditional NR and LS-NR schemes with only a slight increase in computational cost. Importantly, by using an algorithm tailored for constitutive model integration, the current implementation was able to solve the problem of a rod loaded in tension and shear that could not be solved with just a NR implementation and vexed a previous dogleg TR approach. The robustness of the proposed algorithm was studied in detail and demonstrated to exhibit convergence characteristics similar to that of the LS-NR method *when properly scaled* and far in excess of the NR performance. It was also clearly shown that the scaling issues in the TR approach are substantial and the merit function and state variable vector must be scaled to achieve the desired algorithmic performance. Of the two, the merit function was shown to be more potent and the performance of the proposed implementation quickly deteriorates without appropriate treatment.

Given the performance reported through this effort, the proposed routine detailed here represents an exciting possibility towards improved constitutive integration schemes. To this end, the current

implementation is a first attempt at using TR methods that have been developed in the optimization community for other purposes. Potential improvements via specialized model problem formulation or alternatives to the dogleg method could increase efficiency and drive down cost. Additionally, the possibilities of this approach to further improve on the performance for more challenging models with anisotropy, alternative and/or coupled physics (*i.e.* damage or tight thermomechanical coupling), and multisurface cases is enticing. Consideration of cases with alternative inelastic mechanisms and/or physics also necessitates more detailed study of the impact of scaling in these problems. Such cases may introduce additional variables in which a clear choice of scaling constants is not evident due to the differing physical phenomena and unit selection may be unable to mitigate the disparity in variable scaling. These investigations shall be pursued in future efforts.

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