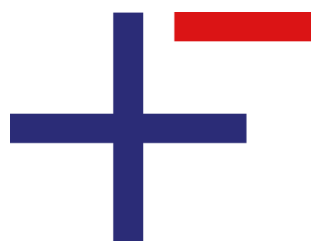


# REDUCED ORDER MODELLING FOR Li-S BATTERIES VIA NUMERICAL SCHEMES

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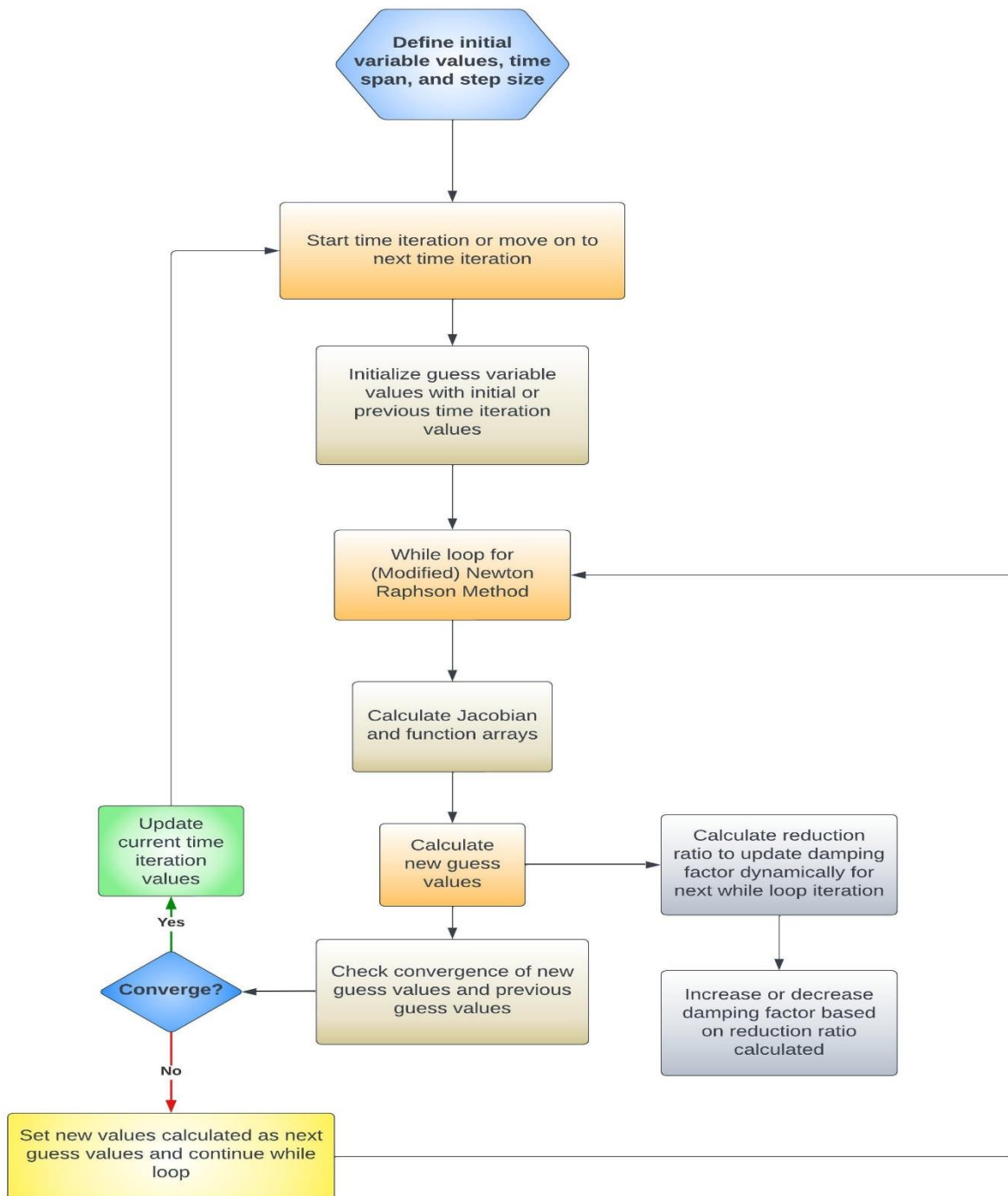
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## 1) BASE MODEL

The base model is coded using a Backwards-Euler discretization and a (modified) Newton Raphson method to solve the system of non-linear equations describing the time evolution of sulfur species defined using the Nernst and Butler-Volmer equations, (1). The base model uses a pre-defined step size which is simple, however less efficient and unstable (takes a time of 10-45 mins to run). The flowchart below shows the outline of how the solver algorithm functions:



The table below highlights the different hyper-parameters used in the solver, their respective definitions and common values used:

Hyper-parameter	Definition	Common values
lamda	Damping factor used to damp (reduce) Jacobian matrix in case of det(Jacobian) approaching large values.	1.0, 0.8 (Starting value, dynamically updated)
damping_update_factor	Used to dynamically update the damping factor (lamda) for different while loop iterations based on ratio of actual reduction to predicted reduction.	0.5, 0.25
damping_min	Minimum value of damping factor	1e-8, 1e-16
regularization_factor	Used as a sort of penalty method, for which the factor is added to the leading diagonal of the Jacobian matrix to ease the instability of the non-linear system of coupled equations.	5e-4, 2.5e-4
n_damp	Used to raise the power of the damping_update_factor for faster or slower damping.	1, 2

The code snippet below shows how these hyper-parameters are used in conjunction of calculating the reductions. NOTE: it is also worth experimenting on trying different values for the highlighted (in red) values which corresponds to the upper and lower bound of the reduction ratio used to update the damping factor:

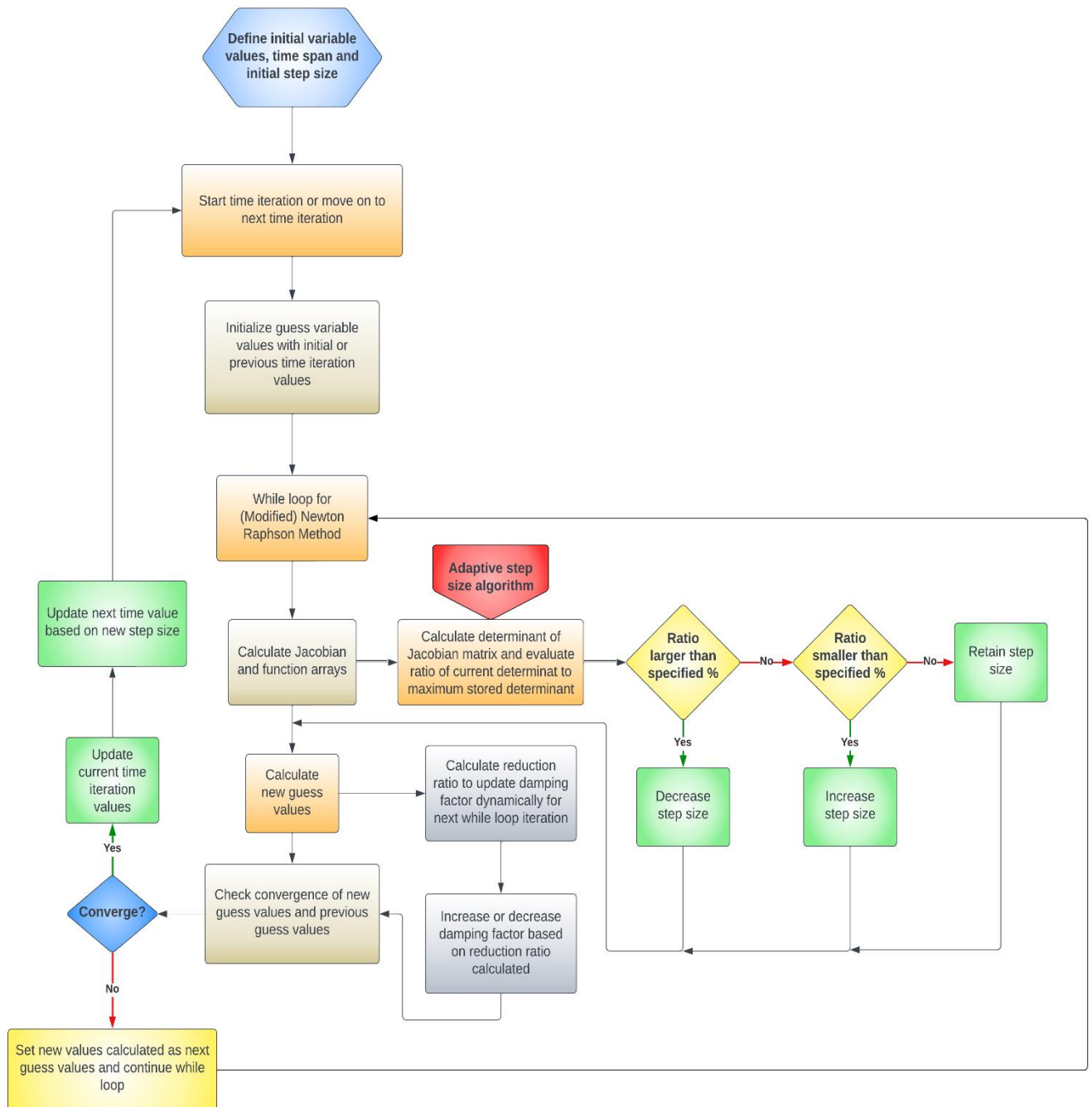
```

388     # Now we solve as usual the new step size will be implemented in the next iteration
389     jacob = jacob + regularization_factor * np.eye(len(x))
390     jacobinv = np.linalg.inv(jacob)
391     # Calculate new values via Newton-Raphson
392     delta = - np.matmul(jacobinv, u_array)
393     new_val = x + lamda*delta
394     new_val = abs(new_val)
395
396     model2 = LiSModel(new_val, I) ## Initialize the model
397     ## Change any values if a new value wants to be used apart from the default ones in the model
398     model2.update_parameters(**upd_params)
399     unew_array = model2.f(h, s8[-1], s4[-1], s2[-1], s[-1], V[-1], sp[-1])
400
401     # Compute the ratio of actual reduction to predicted reduction
402     actual_reduction = np.linalg.norm(u_array) - np.linalg.norm(unew_array)
403     predicted_reduction = np.linalg.norm(u_array) - np.linalg.norm(u_array + jacob @ delta)
404     ratio = abs(actual_reduction / predicted_reduction)
405
406     # Update the damping factor based on the ratio
407     n_damp = 1
408     if ratio > 1e-3:
409         lamda *= (damping_update_factor**n_damp)
410     elif ratio < 1e-4:
411         lamda /= (damping_update_factor**n_damp)
412
413     # Ensure the damping factor does not go below the minimum value
414     lamda = max(lamda, damping_min)
415

```

## 2) ADAPTIVE STEP SIZE MODEL

The next model attempted was the adaptive step size model. This model is an extended version of the base model for which instead of using a fixed step size, the step size is dynamically updated via calculating the ratio of the determinant of the Jacobian for the current iteration to the maximum stored determinant. This model shows a significant increase in both stability and computation time and efficiency (**takes only 2-3s to run**). The flowchart below shows the workflow of the model solver:



As mentioned, this model is a huge improvement to the base model which takes a lengthy time of between 10-45 mins to run a single discharge simulation depending on the step size chosen, however this modified model adaptively updates the step which allows the model to account for unstable solution regions more precisely and increases computational speed for stable solution regions. The table below shows the new added hyper-parameters for the adaptive step size algorithm on top of the previous hyper-parameters:

Hyper-parameters	Definition	Common values
min_h	Minimum allowed step size	1e-4, 1e-6
max_h	Maximum allowed step size	0.5, 1
<code>h_new = max(h*(0.2), min_h)</code>	The red highlighted value is the factor used to reduce the step size by every iteration.	0.05, 0.2, 0.25, 0.5
<code>h_new = min(h/(0.75), max_h)</code>	The red highlighted value is the factor used to increase the step size by every iteration.	0.25, 0.5, 0.75

The code snippet below shows how the adaptive step size algorithm is adapted into the solver.

NOTE: the red highlighted values are also worth experimenting with as they represent the values for the upper bound of Jacobian determinant ratio to decrease the step size (common values: 1.2, 1.5) and lower bound to increase the step size (common values, 0.2, 0.5, 0.75, 1.0):

```

315 # =====
316 #   ### This will dynamically update the step size every iteration ###
317 # =====
318 if i > 1: ## Only check after 1st iteration ##
319     ## Continuously update the maximum value of determinant of Jacobian encountered
320     max_jacobian = max(max_jacobian, abs(jacob_array[i-3]))
321     ## Calculate the ratio of current determinant to maximum for Jacobian
322     max_ratio = abs(jacob_array[i-2])/max_jacobian
323     ## These values need configuration
324     if max_ratio >= 1.5: ## This indicates step needs to be reduced
325         h_new = max(h*(0.25), min_h) ## Saturate at minimum step size
326     elif max_ratio <= 0.2: ## This indicates step size can be increased
327         h_new = min(h/(0.75), max_h) ## Saturate at maximum step size
328     else:
329         h_new = h
330 else:
331     h_new = h

```

### 3) PARAMETER BACKTRACKING MODEL

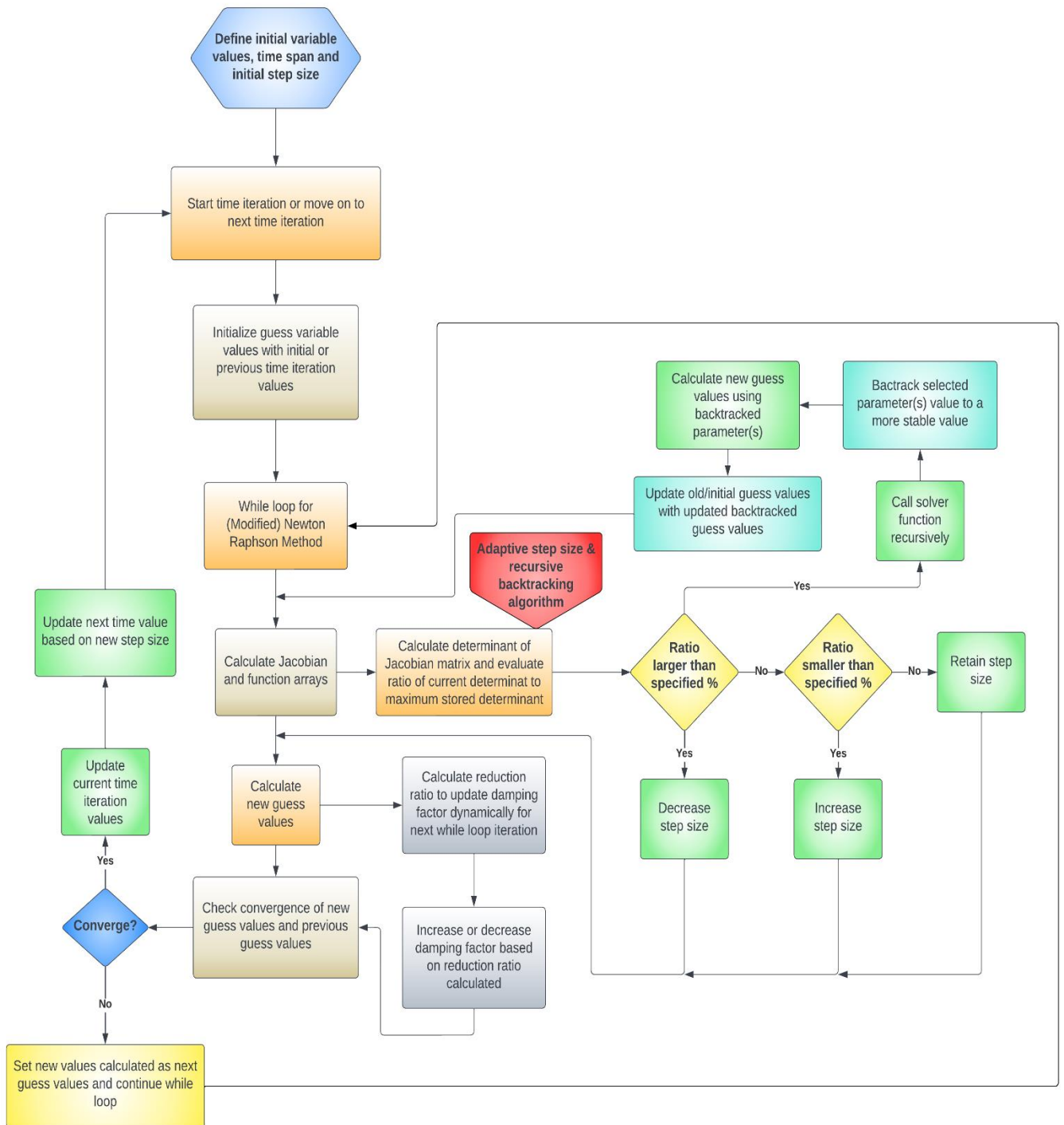
The next model that was devised was the parameter backtracking model, for which the model from the previous adaptive step size is extended to include a backtracking recursive algorithm. This recursive solver call backtracks the solution (at unstable solution regions) to update the guess values with new guesses calculated using stable parameter values, hence labelled backtracking. The code snippet below shows how this new algorithm is used in conjunction with the adaptive step size model:

```
301 # =====
302 #   ### This will dynamically update the step size every iteration ###
303 #   =====
304 if i > 1: ## Only check after 1st iteration ##
305     ## Continuously update the maximum value of determinant of Jacobian encountered
306     max_jacobian = max(max_jacobian, abs(jacob_array[i-3]))
307     ## Calculate the ratio of current determinant to maximum for Jacobian
308     max_ratio = abs(jacob_array[i-2])/max_jacobian
309     ## These values need configuration
310     if max_ratio >= 1.2: ## This indicates step needs to be reduced and parameter backtracking
311         ## Define recursive function call for parameter backtracking
312         ## Recursive call to only solve for 1 iteration:
313         t02 = 0
314         t_end2 = t02 + h
315         new_guess = LiS_Solver(s8guess, s4guess, s2guess, sguess, Vguess, spguess,
316                               t_end2, h, I, break_voltage, state=state, t0=t02, backtracked=True,
317                               params_backtrack=params_backtrack, upd_params=upd_params)
318
319         ## Now update the guess values and run solver by updating u_array and jacobian
320         x_upd = np.array([new_guess[0][-1], new_guess[1][-1], new_guess[2][-1],
321                           new_guess[3][-1], new_guess[4][-1], new_guess[5][-1]])
322         # Update the model
323         model = LiSModel(x_upd, I)
324         ## Change any values if a new value wants to be used apart from the default ones in the model class
325         model.update_parameters(**upd_params)
326         u_array = model.f(h, s8[-1], s4[-1], s2[-1], s[-1], V[-1], sp[-1])
327         jacob = model.jacobian(h)
328         x = x_upd ## Use updated guess values from backtracking
329
330         # Update h (step-size)
331         h_new = max(h*(0.2), min_h) ## Saturate at minimum step size
332
333     elif max_ratio <= 1.0: ## This indicates step size can be increased
334         h_new = min(h/(0.75), max_h) ## Saturate at maximum step size
335
336     else:
337         h_new = h
338
339     else:
340         h_new = h
341
```

NOTE: This snippet is within the LiS\_Solver function, which as can be seen is called recursively. This recursive call is ensured to only occur for a single time step. The backtracked parameter is used to recalculate the guess values and update the current guess values. Through this the model solver seems to achieve greater stability compared to the previous solver iterations.

The flowchart in the next page shows the workflow of the improved solver via backtracking. As it can be seen that all the solver iterations are based on the base model with slight added complexity:

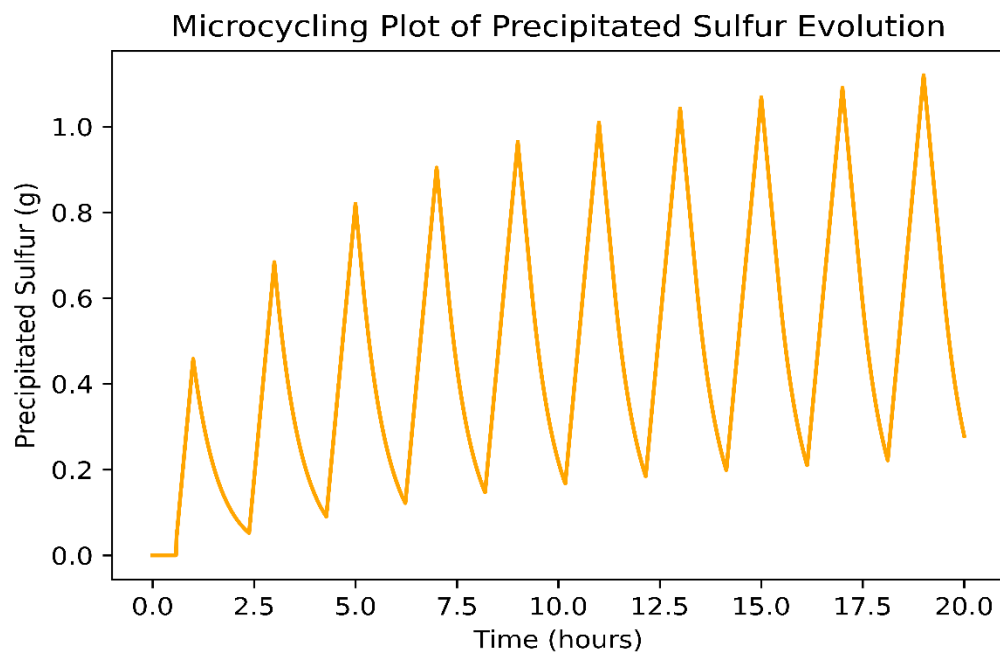
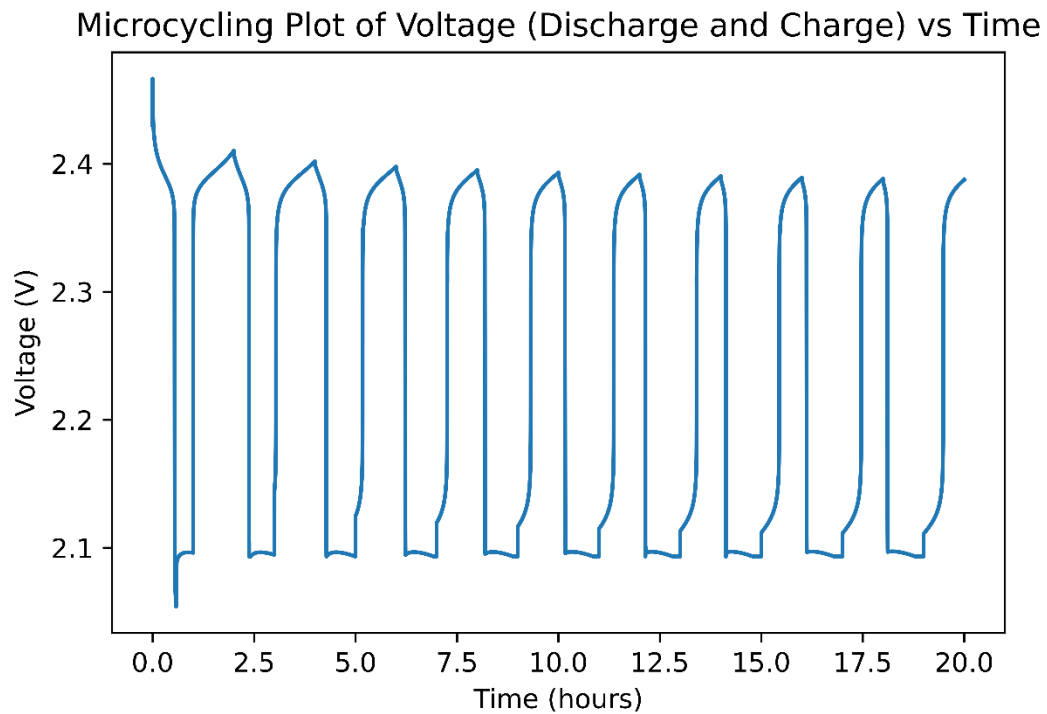




The new algorithm has no newly added hyper-parameters and only uses the previously defined hyper-parameters in the previous sections.



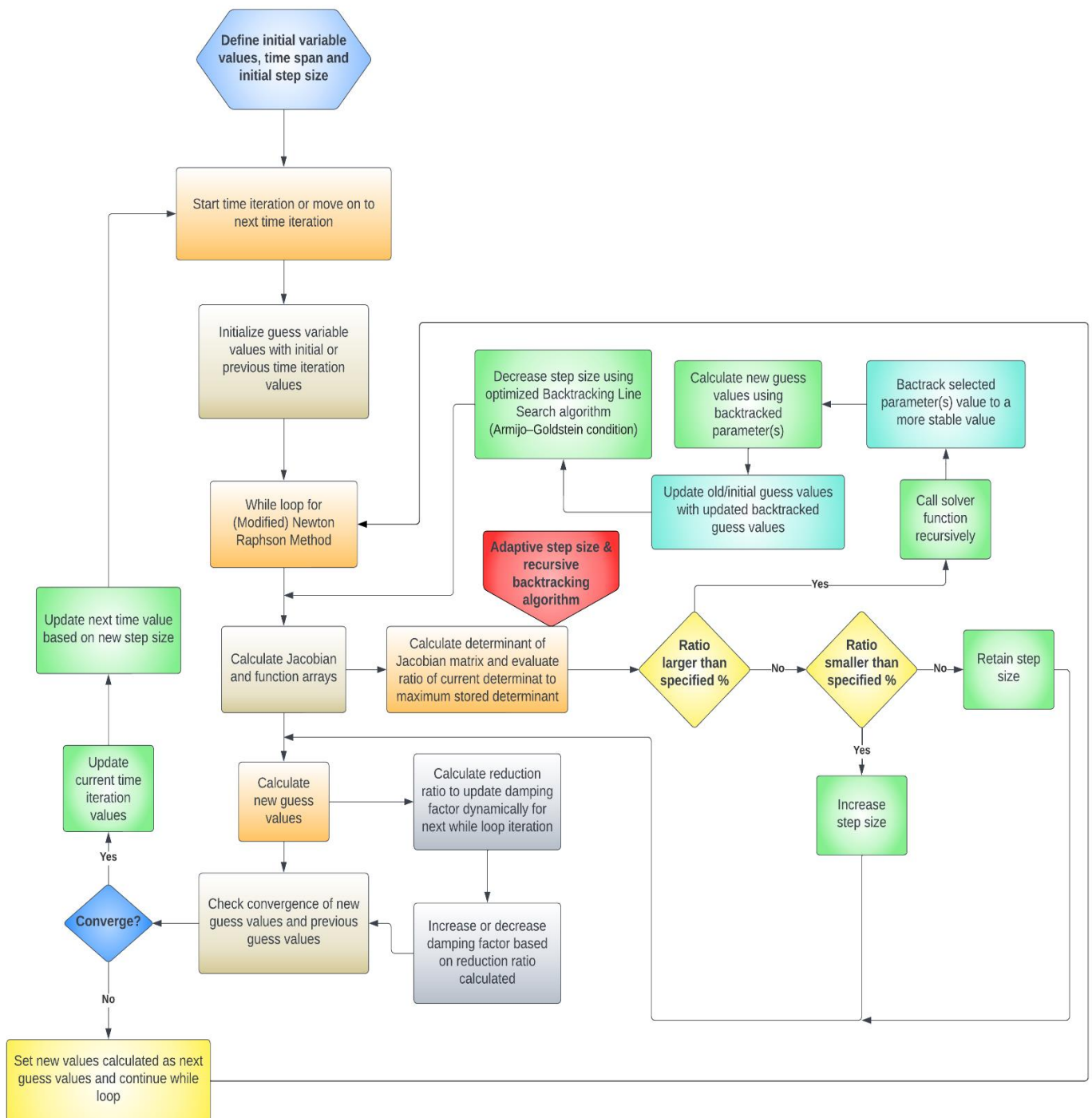
The plots below show the voltage evolution and precipitated sulfur evolution for micro-cycling over 10 cycles (20 hours) using the new parameter backtracking model ( $EH0=2.4$ ,  $EL0=2.0$ ):



#### 4) OPTIMIZED MODEL

The previous backtracking model is further optimized to enable reduction of the step size using the Line-Search optimization method. This method is then used to simulate a new formulation of the system of equations from (2).

The flowchart below is the updated version of the model:



The new optimized model has 2 new hyper-parameters, namely alpha and beta used to represent the Armijo-Goldstein condition. The hyper-parameter and usual values are given in the table below:

Hyper-parameters	Definition	Common values
alpha	Parameter used for Armijo-Goldstein condition	0.25, 0.3, 0.5
beta	Used to decrease step-size	0.05, 0.1, 0.25

The screen-snip below shows the part of the code (within the solver function) in which this Line-Search algorithm is adapted:

```

211 # =====
212 # Define Line Search Method to further optimize the step size
213 # =====
214 jacobinv2 = np.linalg.inv(jacob)
215 delta2 = - np.matmul(jacobinv2,u_array)
216 alpha = 0.3
217 beta = 0.05
218 var_val = x.copy()
219 unew_array = u_array.copy()
220
221 while np.linalg.norm(unew_array) > np.linalg.norm(u_array + alpha*h*np.dot(jacob, delta2)):
222     upd_x = var_val + h*delta2
223     upd_x = abs(upd_x)
224     model3 = LiSModel(upd_x, I) ## Initialize the model
225     ## Change any values if a new value wants to be used apart from the default ones in the model class
226     model3.update_parameters(**upd_params)
227     unew_array = model3.f(h, x_var[:, -1])
228     h *= beta
229     if h < min_h:
230         break
231     #print(h, I*t[i-1]/3600)
232
233     # Further Update h (step-size)
234     h_new = max(h*(0.25), min_h) ## Saturate at minimum step size
235
236     elif max_ratio <= 1.0: ## This indicates step size can be increased
237         h_new = min(h/(0.75), max_h) ## Saturate at maximum step size
238
239     else:
240         h_new = h
241
242     else:
243         h_new = h
244
245     ## Now we solve as usual the new step size will be implemented in the next iteration
246     jacob = jacob + regularization_factor * np.eye(len(x))
247     jacobinv = np.linalg.inv(jacob)
248     # Calculate new values via Newton-Raphson

```

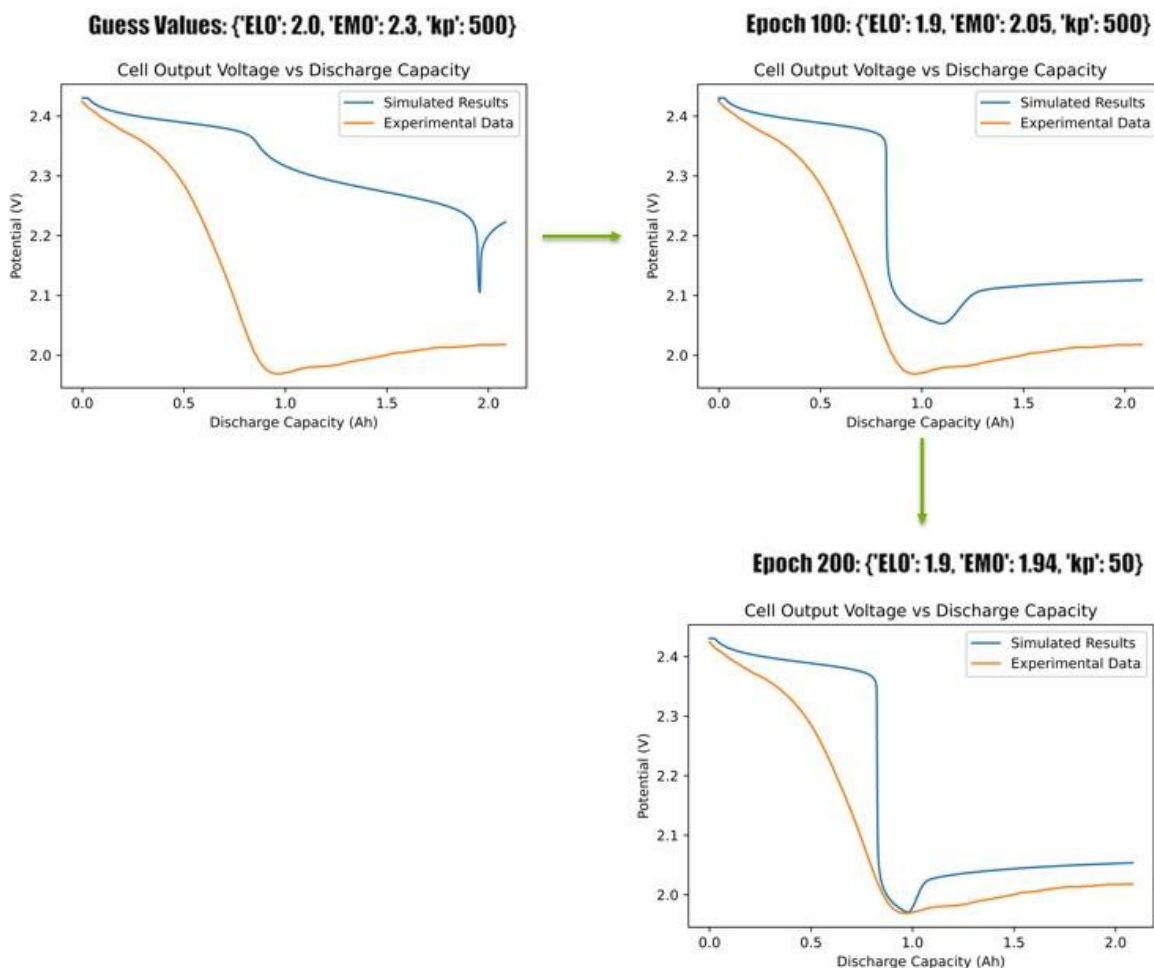
These values for the hyper-parameters, as well as the previous hyper-parameters can be experimented with to find suitable values to ensure the stability of the system that is being solved for.

Finalized Python scripts can be found in the GitHub link below:

[https://github.com/Dharshannan/FUSE\\_Li-S\\_Battery\\_Modelling/tree/main/Finalized\\_Scripts](https://github.com/Dharshannan/FUSE_Li-S_Battery_Modelling/tree/main/Finalized_Scripts)

## 5) GRADIENT DESCENT FOR PARAMETER OPTIMIZATION USING EXPERIMENTAL RESULTS

The new 2022 model formulation (2), is used together with the novel solver within a gradient descent scheme to optimize parameterization using experimental results. The gradient descent scheme is an optimized version of the base gradient descent using Adaptive Moment (ADAM). The result from the gradient descent is as below:



As it can be seen above, the gradient descent starts with a guess value for the 3 optimized parameters here, the low-plateau standard potential (ELO), middle-plateau standard potential (EMO) and the precipitation coefficient (kp). It takes the scheme 200 epochs (\*iterations) to reach optimized parameterization to fit the experimental curve.

More information on how the gradient descent scheme is implemented can be found in the user manual document, which explains how to set up the model and use the solver together with examples for micro-cycling and gradient descent.

## 6) REFERENCES

1. Marinescu M, Zhang T, Offer GJ. A zero dimensional model of lithium-sulfur batteries during charge and discharge. *Physical Chemistry Chemical Physics*. 2016;18(1): 584–593. <https://doi.org/10.1039/c5cp05755h>.
2. Cornish M, Marinescu M. Toward Rigorous Validation of Li-S Battery Models. *Journal of The Electrochemical Society*. 2022;169(6): 060531. <https://doi.org/10.1149/1945-7111/ac7750>.

## APPENDIX

All the models discussed are formulated using the Backward-Euler method, the equations formed are expressed as functions, and these functions are differentiated partially with respect to its variables, the partial differentials are collected to form the Jacobian matrix and the model is then solved using the Newton-Raphson approach. Below are all the functions:

These are an older formulation of the system of ODEs from (1).

$$u1 = h \left( \left( \frac{(n_{s8} \cdot M_{s8} \cdot iH0 \cdot ar)}{n_e \cdot F} \right) \left( \frac{e^{2 \cdot F \cdot (V_{j+1} - EH0) / (R \cdot T)}}{\sqrt{f_h \cdot S_{8,j+1}}} \cdot S_{4,j+1} - \frac{e^{2 \cdot F \cdot (EH0 - V_{j+1}) / (R \cdot T)}}{S_{4,j+1}} \cdot \sqrt{f_h \cdot S_{8,j+1}} \right) - k_s \cdot S_{8,j+1} \right) - S_{8,j+1} + S_{8,j}$$

$$u2 = h \left( \left( -\frac{(n_{s8} \cdot M_{s8} \cdot iH0 \cdot ar)}{n_e \cdot F} \right) \left( \frac{e^{2 \cdot F \cdot (V_{j+1} - EH0) / (R \cdot T)}}{\sqrt{f_h \cdot S_{8,j+1}}} \cdot S_{4,j+1} - \frac{e^{2 \cdot F \cdot (EH0 - V_{j+1}) / (R \cdot T)}}{S_{4,j+1}} \cdot \sqrt{f_h \cdot S_{8,j+1}} \right) + k_s \cdot S_{8,j+1} + \right. \\ \left. \left( \frac{n_{s4} \cdot M_{s8} \cdot iL0 \cdot ar}{n_e \cdot F} \right) \left( \frac{e^{2 \cdot F \cdot (V_{j+1} - EL0) / (R \cdot T)}}{\sqrt{f_l \cdot S_{4,j+1}}} \cdot \sqrt{S_{2,j+1} \cdot (S_{g,j+1})^2} - \frac{e^{2 \cdot F \cdot (EL0 - V_{j+1}) / (R \cdot T)}}{\sqrt{S_{2,j+1} \cdot (S_{g,j+1})^2}} \cdot \sqrt{f_l \cdot S_{4,j+1}} \right) \right) - S_{4,j+1} + S_{4,j}$$

$$u3 = h \left( \left( -\frac{(n_{s2} \cdot M_{s8} \cdot iL0 \cdot ar)}{n_e \cdot F} \right) \left( \frac{e^{2 \cdot F \cdot (V_{j+1} - EL0) / (R \cdot T)}}{\sqrt{f_l \cdot S_{4,j+1}}} \cdot \sqrt{S_{2,j+1} \cdot (S_{g,j+1})^2} - \frac{e^{2 \cdot F \cdot (EL0 - V_{j+1}) / (R \cdot T)}}{\sqrt{S_{2,j+1} \cdot (S_{g,j+1})^2}} \cdot \sqrt{f_l \cdot S_{4,j+1}} \right) \right) - S_{2,j+1} + S_{2,j}$$

$$u4 = h \left( \left( -2 \cdot \frac{(n_s \cdot M_{s8} \cdot iL0 \cdot ar)}{n_e \cdot F} \right) \left( \frac{e^{2 \cdot F \cdot (V_{j+1} - EL0) / (R \cdot T)}}{\sqrt{f_l \cdot S_{4,j+1}}} \cdot \sqrt{S_{2,j+1} \cdot (S_{j+1})^2} - \frac{e^{2 \cdot F \cdot (EL0 - V_{j+1}) / (R \cdot T)}}{\sqrt{S_{2,j+1} \cdot (S_{j+1})^2}} \cdot \sqrt{f_l \cdot S_{4,j+1}} \right) - \left( \frac{k_p \cdot S_{p,j+1}}{v \cdot \rho_s} \cdot (S_{j+1} - S_{sat}) \right) \right) - S_{j+1} + S_j$$

$$u5 = I + iH0 \cdot ar \left( \frac{e^{2 \cdot F \cdot (V_{j+1} - EH0) / (R \cdot T)}}{\sqrt{f_h \cdot S_{8,j+1}}} \cdot S_{4,j+1} - \frac{e^{2 \cdot F \cdot (EH0 - V_{j+1}) / (R \cdot T)}}{S_{4,j+1}} \cdot \sqrt{f_h \cdot S_{8,j+1}} \right) + iL0 \cdot ar \left( \frac{e^{2 \cdot F \cdot (V_{j+1} - EL0) / (R \cdot T)}}{\sqrt{f_l \cdot S_{4,j+1}}} \cdot \sqrt{S_{2,j+1} \cdot (S_{j+1})^2} - \frac{e^{2 \cdot F \cdot (EL0 - V_{j+1}) / (R \cdot T)}}{\sqrt{S_{2,j+1} \cdot (S_{j+1})^2}} \cdot \sqrt{f_l \cdot S_{4,j+1}} \right)$$

$$u6 = \frac{h \cdot k_p \cdot S_{p,j+1}}{v \cdot \rho_s} \cdot (S_{j+1} - S_{sat}) - S_{p,j+1} + S_{p,j}$$

The equations are partially differentiated with respect to each of the (j+1) variables, the Jacobian is obtained and model solved using Newton-Raphson, as detailed below:

$$\begin{bmatrix} S_{8,j+1} \\ S_{4,j+1} \\ S_{2,j+1} \\ S_{j+1} \\ V_{j+1} \\ S_{p,j+1} \end{bmatrix}_{(n+1)} = \begin{bmatrix} S_{8,j+1} \\ S_{4,j+1} \\ S_{2,j+1} \\ S_{j+1} \\ V_{j+1} \\ S_{p,j+1} \end{bmatrix}_{(n)} - \begin{bmatrix} \frac{\partial u1}{\partial s_{8,j+1}} & \frac{\partial u1}{\partial s_{4,j+1}} & 0 & 0 & \frac{\partial u1}{\partial V_{j+1}} & 0 \\ \frac{\partial u2}{\partial s_{8,j+1}} & \frac{\partial u2}{\partial s_{4,j+1}} & \frac{\partial u2}{\partial s_{2,j+1}} & \frac{\partial u2}{\partial s_{j+1}} & \frac{\partial u2}{\partial V_{j+1}} & 0 \\ 0 & \frac{\partial u3}{\partial s_{4,j+1}} & \frac{\partial u3}{\partial s_{2,j+1}} & \frac{\partial u3}{\partial s_{j+1}} & \frac{\partial u3}{\partial V_{j+1}} & 0 \\ 0 & \frac{\partial u4}{\partial s_{4,j+1}} & \frac{\partial u4}{\partial s_{2,j+1}} & \frac{\partial u4}{\partial s_{j+1}} & \frac{\partial u4}{\partial V_{j+1}} & \frac{\partial u4}{\partial s_{p,j+1}} \\ \frac{\partial u5}{\partial s_{8,j+1}} & \frac{\partial u5}{\partial s_{4,j+1}} & \frac{\partial u5}{\partial s_{2,j+1}} & \frac{\partial u5}{\partial s_{j+1}} & \frac{\partial u5}{\partial V_{j+1}} & 0 \\ 0 & 0 & 0 & \frac{\partial u6}{\partial s_{j+1}} & 0 & \frac{\partial u6}{\partial s_{p,j+1}} \end{bmatrix}_{(n)}^{-1} \cdot \begin{bmatrix} u1 \\ u2 \\ u3 \\ u4 \\ u5 \\ u6 \end{bmatrix}_{(n)}$$

The (n) terms represent the n<sup>th</sup> guess values and the (n+1) terms represents the next guess values or new values if converged.

The method to change from the older 2016 model formulation (1) to the newer 2022 formulation (2), follows the same manner as depicted above for the older formulation.