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A Statistical Test for Identifying the Number of Creep Regimes when using the Wilshire Equations for Creep Property Predictions

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9 ABSTRACT

10 A new parametric approach, termed the Wilshire equations, offer the realistic potential of being able to accurately life materials operating at in service conditions from accelerated test results 11 lasting no more than 5,000h. The success of this approach can be attributable to a well-defined 12 linear relationship that appears to exist between various creep properties and a log 13 14 transformation of the normalised stress. However, these linear trends are subject to 15 discontinuities, the number of which appears to differ from material to material. These discontinuities have until now been i. treated as abrupt in nature and ii. been identified by eye 16 17 from an inspection of simple graphical plots of the data. This paper puts forward a statistical test for determining the correct number of discontinuities present within a creep data set and a 18 19 method for allowing these discontinuities to occur more gradually - so that the methodology is more in line with the accepted view as to how creep mechanism evolve with changing test 20 conditions. These two developments are fully illustrated using creep data sets on two steel 21 alloys. When these new procedures are applied to these steel alloys, not only do they produce 22 23 more accurate and realistic looking long-term predictions of the minimum creep rate but they 24 also lead to very different conclusions about the mechanisms determining the rates of creep from those originally put forward by Wilshire. 25

26 Keywords:

27

- 28
- 29 30

I. INTRODUCTION

Minimum creep rate, Wilshire methodology, Statistical testing, Threshold models

To reduce fuel consumption and CO2 emissions from power plants, new high-temperature 31 32 alloys are required to resist the increase in temperature and pressure needed to raise plant efficiencies. However, at the design stage, information must be available on the stresses to 33 which multiple batches of these new alloys can sustain without creep fracture occurring within 34 35 100,000h at the service temperatures ^[1]. Unfortunately, with the traditional parametric, numerical and computational methods, long term strengths cannot be predicted by 36 extrapolation of short-term property sets. Consequently, at present, protracted and expensive 37 long-duration test programmes are necessary to determine the 100,000h creep rupture 38

strengths, with a reduction in this 12 to 15 year "materials development cycle" being defined
 as the No.1 priority in the 2007 UK Energy Materials-Strategic Research ^[2].

In response to this problem, over recent years, a new approach - termed the Wilshire 41 42 equations - has been devised which appears to allow accurate long-term strength values to be obtained by extrapolation from accelerated short-term measurements. The last 5 to 6 years has 43 seen the appearance in the literature of this methodology applied to a wide range of materials 44 used for high temperature application in the power generation and aerospace industries in an 45 attempt to verify the validity and accuracy of this approach [3-8]. Specifically, 100,000h strength 46 estimates have been produced by analysis of multi-batch data lasting up to only 5,000h for a 47 series of ferritic bainitic and martensitic steels for power and petrochemical plant and titanium 48 49 alloys used in aero engine blades and disc.

50 The Wilshire equation takes the form,

51
$$\left(\sigma/\sigma_{TS}\right) = \exp\left\{-k_2\left[\dot{\varepsilon}_m \cdot \exp(Q_c^*/RT)\right]^v\right\}$$
 [1a]

where $\dot{\epsilon}_{m}$ is the minimum creep rate, T is the absolute temperature, σ the stress, σ_{TS} the tensile strength, R the universal gas constant, Q^{*}_{c} the activation energy for self-diffusion and where k_{2} and v are further model parameters. This equation provides a sigmoidal data presentation such that $\dot{\epsilon}_{m} \rightarrow \infty$ as $(\sigma/\sigma_{TS}) \rightarrow 1$ (provided v < 0), whereas $\dot{\epsilon}_{m} \rightarrow 0$ as $(\sigma/\sigma_{TS}) \rightarrow 0$. Wilshire and Battenbough ^[3] proposed a very similar expression to Eq. [1] for the stress and temperature dependencies of the time to failure, t_{f} , and time to various different strains. The parameters k_{2} and v appear to be dependent upon stress (and possibly temperature) for many steel alloys.

59 This approach can be contrasted to the traditional power law expression for modelling 60 creep properties as a function of stress and temperature

61
$$\dot{\varepsilon}_{\rm m} = A\sigma^n \exp(Q_{\rm c}^*/{\rm RT})$$
 [1b]

but once again the unknown parameter $(Q_c^* and n)$ change with test conditions. In this approach 62 the variation in n and O_{c}^{*} with test conditions is traditionally explained in terms of differing 63 creep mechanisms being dominant at different stresses and temperatures. For example, a 64 transition from $n \approx 4$ to $n \approx 1$ is traditionally taken as evidence of a change from dislocation to 65 66 diffusional creep processes as stress diminishes. Likewise, when creep occurs by diffusion controlled generation and movement of dislocations a fall in the activation energy below that 67 68 associated with lattice self-diffusion is interpreted either as i. deformation behaviour being 69 increasingly controlled by preferential diffusion along dislocation cores at low temperatures 70 within a high stress regime, or by ii. deformation behaviour being increasingly controlled by 71 stress directed vacancy flow along grain boundaries at low temperatures and stresses.

However, the results obtained from using Eq. [1a] have lead authors like Wilshire and Scharning ^[4] and Wilshire and Whittaker ^[5] to suggest that the parameter instability observed in k_2 and v is not the result of a change from dislocation to diffusional creep processes. Instead, and depending on the material under investigation, they choose to interpret the observed changes in k_2 and v as either being: i. the result of particle coarsening associated with long test durations at lower stresses.
ii. or as a result of a change from creep occurring from the generation of new dislocations
within the lattice structure itself to creep occurring from the movement of dislocations preexisting only in the grain boundary zones as a result of a low stress level.

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The fact that Eq. [1a] has been remarkably successful in being able to predict creep key lives at operating conditions from highly accelerated tests of very short duration and over a wide range of materials is taken by these authors to be strong evidence to support this view.

As an illustration of this point of view consider two steel alloys. Fig.1a summarises the 86 results obtained by Wilshire and Scharning^[4] in their 2007 study of 1Cr - 1Mo - 0.25V steel 87 using the NIMS^[9] data base on this material. As can be seen from this figure, there appears to 88 be one break point (and therefore two creep regimes) where the values for k₂ and v change, but 89 according to the authors, the activation energy remains unchanged. By studying the 90 metallographic evidence obtained by NIMS^[10], the authors found that little or no change was 91 92 observed in the as received bainitic microstructures when hardness reductions were small, 93 whereas distinguishable increases in carbide size was apparent when the hardness values fell of rapidly. Furthermore, only very modest falls in hardness were observed in the high 94 normalised stress range, with rapid harness reductions occurring the low normalised stress 95 96 ranges. Thus, the unchanging activation energy quoted by the authors is taken to mean that creep is determined by behaviour within the crystal lattice. Then the changes in k₂ and v reflect 97 98 differences in the rates of creep strength reduction caused by the evolution of the tempered bianitic microstructure in the low normalised stress range. This causes creep rates to be much 99 higher in the low stress regime than would be predicted by relations prevalent at higher stresses. 100

101 Thus in Fig.1a the larger carbide particle sizes present at very low stresses (where the 102 test duration is long), means that at a given stress, creep rates will be greater than that predicted 103 from relations that hold at higher stresses. Hence the steeper slope of the best fit line shown in 104 Fig.1a below a normalised stress of around 0.4. Despite this, and as clearly seen in Fig. 1a, the 105 presence of these distinctly different stress regimes does not prevent the accurate prediction of 106 creep lives out to over 100,000 hours using only data up to 5,000h for the purpose of parameter 107 estimation.

In their study of 2.25Cr-1Mo steel, Wilshire and Whittaker ^[5] identified three different 108 values for v and k_2 that corresponded to high, medium and low stress regimes – as seen in Fig. 109 1b for the MAF batch of materials within the NIMS^[11] data base on this steel alloy. For this 110 material, these authors again suggest that no transition takes place from dislocation to 111 diffusional creep with decreasing applied stress. Instead, dislocation creep processes are rate 112 controlling at all stress levels, even though the detailed dislocation processes vary in different 113 stress regimes. Thus, with 2.25Cr-1Mo steels, the creep and creep fracture properties differ 114 above and below $\sigma \approx \sigma_Y$ (where σ_Y is the yield stress). According to Wilshire and Whittaker 115 ^[5], when $\sigma > \sigma_{\rm Y}$, so that the initial strain on loading has both elastic and plastic components, 116 117 creep is controlled by the generation and movement of dislocations within the grains where the 118 activation energy is highest.

In contrast, when $\sigma < \sigma_{\rm Y}$, so that the strain on loading has essentially only an elastic 119 component, new dislocations are not generated within the grains. Instead, creep occurs within 120 the grain boundary zones, i.e. by grain boundary sliding or diffusion along existing dislocations 121 and grain boundaries with associated deformation in the grain regions adjacent to the 122 boundaries (where the activation energy is lower). Hence, the creep rates when $\sigma < \sigma_{\rm Y}$ are 123 slower and the creep lives are longer than expected by direct extrapolation of $\dot{\mathcal{E}}_m$ data obtained 124 125 when $\sigma < \sigma_{Y}$. Another change in creep and creep rupture behaviour occurs when σ 126 approximately equals $0.2\sigma_{TS}$. With this material, the original ferrite/bainite microstructure 127 degrades to ferrite and molybdenum carbide particles in long term tests at the highest creep temperatures, with very coarse carbide particles forming along the grain boundaries. This 128 carbide coarsening reduces creep strength in the matrix allowing diffusion to occur within the 129 grains once again where the activation energy is higher. In these cases, because of the loss of 130 creep resistance caused by this transformation, the $\dot{\mathcal{E}}_m$ values are larger when $\sigma < 0.2\sigma_{\rm TS}$ than 131 would be predicted by extrapolation of data collected at intermediate σ levels. These authors 132 133 have provided similar explanations for the observed breaks in other power generating materials as well. 134

Yet despite the simplicity of these types of explanation, and the accuracy of predictions 135 of creep life made using this approach, the methodology has always been presented (with Figs 136 .1a,b being a typical visualisation of the approach in the literature) showing an abrupt change 137 in parameter values at precise values for the normalised stress. This suggests that at this 138 139 normalised stress the cause of creep deformation suddenly changes from being 100% controlled by dislocations within the bulk to 100% determined by dislocations within the 140 boundaries. Yet, such changes are known to occur gradually, with a gradual transition say from 141 142 deformation being controlled by the bulk to being controlled within the grains as stress falls below a critical value. 143

144 This paper therefore has two main aims designed to enhance and further formulise the Wilshire methodology. The first is to modify the Wilshire methodology to allow for a gradual 145 rather than abrupt change by using the approach first put forward by Evans^[12] - but to generalise 146 this approach to allow for more than one "regime" change. Secondly, a statistical test is 147 presented that enables the number of regime changes or breaks present in the creep data to be 148 149 determined. Such a statistical test is not as straight forward as it first sounds because under the null hypothesis of no regime change some of the parameters in the Wilshire equations are not 150 actually defined. As a consequence of this, the distribution of any test statistic for this null 151 hypothesis is non-standard - as maximum likelihood (or least squares) theory is no longer 152 153 directly applicable. Interestingly, the modified Wilshire methodology proposed here provides 154 a neat solution to this problem of testing for regime change.

155

II. THE MODIFIED WILSHIRE EQUATIONS

156 A. Two Competing Creep Deformation Mechanisms

157 To develop the proposed modification of the Wilshire equations, it is first helpful to 158 rewrite Eq. [1a] in the following way

159
$$y = a + bx_1 + dx_2 + u_1$$
 [2]

with y being the natural log of the minimum creep rate, $x_1 = \ln[-\ln[\sigma/\sigma_{TS})]$ and $x_2 = 1/RT$, b = 1/v, $a = \ln[k_2/v]$ and $d = Q^*_c$. u_1 are the residuals included in the specification to make clear the fact that the experimental data on creep properties are stochastic in nature. Estimation procedures for determining values for a, b and d typical take the form of minimising the sum of these squared residuals. Consider next the simplest scenario where the data has at most just a single break or two distinct creep mechanisms or regimes. In such a situation Eq. [2] can be written as

167
$$y = \begin{cases} a_1 + b_1 x_1 + d_1 x_2 + u_2 \\ a_2 + b_2 x_1 + d_2 x_2 + u_2 \end{cases}$$
 with proportion 1-w (3a)

where, for example, b_1 is the value for b under one creep mechanism and b_2 the value for b 168 under the other creep mechanism. u_2 are the residuals associated with the Wilshire model that 169 has two creep regimes. The value for w determines how much of the overall minimum creep 170 171 rate is determined by a particular mechanism. So when w = 0.5 two different creep processes 172 (for example dislocation movements within grain boundaries versus dislocation movements 173 within the bulk) contribute equally towards the overall minimum creep rate. Then as w tends to unity (and so 1 - w₁ tends to zero) the creep rate is increasingly determined by just one of 174 175 these creep mechanisms. When w = 1, the creep rate is determined 100% by a single mechanism. In effect w measures the dominance of a particular deformation mechanism. Then 176 d_1 can be interpreted as the activation energy associated with the first mechanism, and d_2 is the 177 activation energy associated with the other mechanism (for example the activation energies 178 associated with dislocation movements within boundaries and within the bulk). In comparison 179 to Eq. [1], $b_1 = 1/v_1$ and $b_2 = 1/v_2$ where v_1 and v_2 are the values for v in Eq. [1] associated with 180 the two different regimes. Likewise, $a_1 = \ln[k_{21}/v_1]$ and $a_2 = \ln[k_{22}/v_2]$ where k_{21} and k_{22} are 181 the values for k_2 in Eq [1] associated with the two different regimes. 182

183 Whilst it is unclear exactly how w varies with the normalised stress, it must be the case 184 that w tends 1 as σ/σ_{TS} increases. Whilst this could happen in a linear fashion, a more general 185 representation would allow for a non-linear transition between the regimes

186
$$\mathbf{w} = \frac{1}{1 + \exp[-\beta_1(\mathbf{x}_1 - \mathbf{x}_1^*)]}$$
 [3b]

187 where x_{1}^{*} is some critical value for the normalised stress, namely that normalised stress where 188 creep rates are equally governed by the two competing mechanisms (i.e. where w = 0.5). The 189 specification given by Eq. [3a,b] is very similar to threshold models used quite commonly for 190 modelling time series data and the reader is referred to Tong ^[12] and Martin et. al. ^[13] for a good 191 review on how to identify and estimate the parameters of such models. Writing the 192 determination of w in this way has the clear advantage that the traditional Wilshire equation 193 can be recovered from this re-specification. That is, if β_1 is large (typically larger than 500), 194 the S shaped sigmoidal curve given by Eq. [3b] becomes extremely steep around x^*_1 and 195 essentially appears as a step like function at this point leading to a very abrupt regime change 196 – which is how the Wilshire equations has been applied up until now. That is, as β_1 increase, 197 Eq. [3b] approximates to the step function

198
$$W = \begin{cases} 1 & \text{if } x_1 \ge x_1^* \\ 0 & \text{if } x_1 < x_1^* \end{cases}$$
 [3c]

However, the main advantage if Eq. [3b] is that unlike a step function implied by the traditional Wilshire model, w is differentiable and this provides a means for statistically testing whether such a regime change exists in the first place.

Eqs. [3a,b] or Eq. [3a,c] can be combined into a single equation of the form

203
$$y = (a_1 + b_1x_1 + d_1x_2 + u_2)(1 - w) + (a_2 + b_2x_1 + d_2x_2 + u_2)w$$

204 or

205 $y = a_1 + b_1 x_1 + d_1 x_2 + (a_2 - a_1)w + (b_2 - b_1)w x_1 + (d_2 - d_1)w x_2 + u_2$ [3d]

206

When the model is expressed as in Eq. [3d], a simple estimation procedure for the 207 unknown parameters can be used. First, arbitrarily choose a value for β_1 and x^*_1 in Eq. [3b]. 208 This makes wan observable variable in Eq. [3d] so that the parameters of this equation can be 209 obtained by regressing y on a constant, x_1 , x_2 , w and the cross products wx_1 and wx_2 (this is 210 just multiple linear least squares the value for u^2_2 summed over all data points is minimised). 211 Then a grid search can be carried out to find the values for β_1 and x^*_1 that further minimise the 212 residual sum of squares (x_1^*) will typically be varied in small increments over the range 0.2 to 213 0.8, whilst β_1 will typically be varied in less small increments over the range 0 to 1000). This 214 will produce estimates for a_1 , b_1 and d_1 together with a_2 - a_1 , b_2 - b_1 and d_2 - d_1 . From all these 215 estimates, it is then possible to recover the values for a_2 , b_2 and d_2 . 216

217 B. Three or More Competing Creep Deformation Mechanisms

There are a number of ways to generalise Eq. [3d]. One is to allow for mechanism changes at more than one normalised stress level - as suggested by Wilshire and Whittaker when studying data on 2.25Cr-1Mo steel. The second is to allow the mechanism to change at various stress and temperature levels as is typically portrayed in traditional creep deformation maps. Such an approach was considered by Evans^[14] and will not be discussed further in this paper. In the former approach, Eq. [3a] would generalise to

$$a_{1} + b_{1}x_{1} + d_{1}x_{2} + u_{3}$$
with proportion w₁

$$a_{2} + b_{2}x_{1} + d_{2}x_{2} + u_{3}$$
with proportion w₂

$$y = a_{3} + b_{3}x_{1} + d_{3}x_{2} + u_{3}$$
with proportion w₃

$$\vdots$$

$$a_{p} + b_{p}x_{1} + d_{p}x_{2} + u_{3}$$
with proportion w_p = 1 - w₁ - w₂ - ... - w_{p-1}
225 [4a]

where u_3 are the residuals when there are p different creep mechanisms that predominantly come into operation at p - 1 different normalised stresses. For example, consider three possible creep regimes that occur within different normalised stress ranges. The Eq. [4a] simplifies to

229
$$y = (a_1 + b_1x_1 + d_1x_2 + u_3)(w_1) + (a_2 + b_2x_1 + d_2x_2 + u_3)(1 - w_1 - w_3) + (a_3 + b_3x_1 + d_3x_2 + u_3)w_3$$

231
$$y = a_2 + b_2 x_1 + d_2 x_2 + (a_1 - a_2) w_1 + (a_3 - a_2) w_3 + (b_1 - b_2) w_1 x_1 + (b_3 - b_2) w_3 x_1 + (d_1 - d_2) w_1 x_2 + (d_3 - d_2) w_3 x_2 + u_3$$
[4b]

233
$$\mathbf{w}_1 = 1 - \frac{1}{1 + \exp[-\beta_1(\mathbf{x}_1 - \mathbf{x}_1^*)]}; \quad \mathbf{w}_3 = \frac{1}{1 + \exp[-\beta_3(\mathbf{x}_1 - \mathbf{x}_1^{**})]}; \quad \mathbf{w}_2 = 1 - \mathbf{w}_1 - \mathbf{w}_3$$
 [4c]

and with x^{**_1} being the normalised stress associated with another creep mechanism starting to dominated the process of deformation. Eq. [4c] allows the intermediate regime to phase in as the regimes either side start to take on a less dominant role.

237 III. A STATISTICAL TEST FOR REGIME CHANGE

238 A. Two Competing Creep Deformation Mechanisms

By testing jointly that the parameters $(a_2 - a_1)$, $(b_2 - b_1)$ and $(d_2 - d_1)$ are all equal to zero 239 in Eq. [3d] it becomes possible to determine statistically how many regime changes are present 240 within the experimental data. A natural test statistic to determine whether or not these are zero 241 (which is the null hypothesis) is to jointly test whether the parameters in front of w, wx_1 and 242 wx₂ in Eq. [3d] are significantly different from zero. (Readers are referred to Vining and 243 Kowalski ^[15] for a description on this joint test of significance). However, the Standard F test 244 normally constructed to carry out such a test, no longer has an F distribution because the 245 parameters β_1 and x_1^* in Eq. [3b] are not defined under this null hypothesis and so conventional 246 maximum likelihood theory is no longer directly applicable. An alternative approach is to test 247 $\beta_1 = 0$ as w is Eq. [3b] then becomes a constant resulting in Eq. [3a] collapsing to Eq. [1]. 248 However, in this case it is the parameters x_1^* , a_1 , b_1 , d_1 , a_2 , b_2 and d_2 that are not identified under 249 the null hypothesis. In the Econometrics literature, three possible ways to address this problem 250 have been identified. Luukkonen, Saikkonen and Terasvirta ^[16] suggest focusing on the local 251 asymptotics at $\beta_1 = 0$. This approach has the advantage of yielding a test statistic with a standard 252 distribution under the null hypothesis. Alternatively, Hansen ^[17] proposes a solution based on 253

local asymptotics at $a_1 = b_1 = d_1 = a_2 = b_2 = d_2 = 0$, which yields a test statistic whose distribution must be approximated by bootstrapping. Lee, Granger and White ^[18] proposes a test similar to Hansen's in that it tests $a_1 = b_1 = d_1 = a_2 = b_2 = d_2 = 0$. However, it draws simulated values for x^*_1 and β_1 to generate values for w in Eq. [3b]. The authors suggest using a rectangular distribution to do this simulation.

This paper makes use of the first of these approaches. Let $z = \beta_1(x_1 - x^*_1)$ so that under the null hypothesis of no regime change $\beta_1 = 0$, and so z = 0. The first three derivatives of Eq. [3b] with respect to z, evaluated at z = 0 are as follows:

262
$$\mathbf{w}_{0}^{(1)} = \frac{\partial w}{\partial z}\Big|_{z=0} = \frac{\exp(-z)}{(1+\exp(-z))^{2}}\Big|_{z=0} = \frac{1}{4}$$

263
$$\mathbf{w}_{0}^{(2)} = \frac{\partial^{2} w}{\partial z^{2}} \bigg|_{z=0} = \frac{\exp(-z) - \exp(-2z)}{(1 + \exp(-z))^{3}} \bigg|_{z=0} = 0$$

264
$$w_{0}^{(3)} = \frac{\partial^{3} w}{\partial z^{3}} \bigg|_{z=0} = \frac{\exp(3z) - 4\exp(2z) + \exp(z)}{6\exp(2z) + 4\exp(3z) + \exp(4z) + 4\exp(z) + 1} \bigg|_{z=0} = -\frac{1}{8}$$

Using these derivatives in a third order Taylor series expansion of w around z = 0 gives

266
$$\mathbf{w} \approx w_{(0)} + w_0^{(1)}(z-0) + \frac{1}{2}w_0^{(2)}(z-0)^2 + \frac{1}{6}w_0^{(3)}(z-0)^3 = \frac{1}{2} + \frac{1}{4}z + 0 - \frac{1}{48}z^3 \quad [5a]$$

Now the expansion of z^3 in Eq. [5a] has terms x_1 , x^2_1 and z^3_1 so that this function can be approximated by the cubic

269 $W \approx \delta_0 + \delta_1 x_1 + \delta_2 x_1^2 + \delta_3 x_1^3$ [5b]

This Taylor series approximation represents the local behaviour of the function in the vicinity of $\beta_1 = 0$ and therefore provides a basis for a test of regime change. Substituting Eq. [5b] into Eq. [3d], (and ignoring the residual term for the moment), gives a regression equation of the form

$$y = \{a_{1} + (a_{2} - a_{1})\delta_{0}\} + \{b_{1} + (a_{2} - a_{1})\delta_{1} + (b_{2} - b_{1})\delta_{0}\}x_{1} + \{d_{1} + (d_{2} - d_{1})\delta_{0}\}x_{2}$$

$$+ \{(a_{2} - a_{1})\delta_{2} + (b_{2} - b_{1})\delta_{1}\}x_{1}^{2} + \{(a_{2} - a_{1})\delta_{3} + (b_{2} - b_{1})\delta_{2}\}x_{1}^{3} + (b_{2} - b_{1})\delta_{3}x_{1}^{4}$$
 [5c]

$$+ (d_{2} - d_{1})\delta_{1}x_{1}x_{2} + (d_{2} - d_{1})\delta_{2}x_{1}^{2}x_{2} + (d_{2} - d_{1})\delta_{3}x_{1}^{3}x_{2}$$

Under the null hypothesis of no regime change, there are no interaction terms and no quadratic, cubic or fourth order terms (as the values in round brackets are then zero) present in Eq. [5c]. Therefore, the steps required to perform a test of the null hypothesis that there is just one creep mechanism (i.e. no creep regime change) are as follows:

Step 1: Regress y on $\{1, x_1, x_2\}$ (i.e. assuming no regime change as in Eq. [2]) to get estimates of the residuals u_1 shown in Eq.[2]. 281 Step 2: Regress u_1 on $\{1, x_1, x_2, x^{2_1}, x^{3_1}, x^{4_1}, x_1x_2, x^{2_1}x_2, x^{3_1}x_2\}$.

Step 3: Compute the Lagrange multiplier statistic $LM = NR^2$ where N is the sample size and R² is the coefficient of determination from the regression carried out in step 2. Under the null hypothesis that $\beta_1 = 0$ (i.e. no regime change), LM is asymptotically distributed as a chi square variable with 6 degrees of freedom.

The intuition behind this test is that any important regime change excluded from the regression in step 1 will show up in the regression carried out in step 2 in the form of a high value for the coefficient of determination R^2 (and so lead to a large chi square variable and the subsequent rejection of the null hypothesis).

290 B. Three or More Competing Creep Deformation Mechanisms

291 This test is easily generalised to three or more competing mechanism by adopting a sequential estimation and testing procedure. Thus, the initial null hypothesis is for a linear 292 model with a single creep mechanism and this is tested against the alternative of a model with 293 294 a single regime change (or two mechanisms) using exactly the same procedure as that outlined in sub section IIIA above. If the null hypothesis is accepted that is the end of this sequential 295 procedure and there is just a single creep mechanism present within the data. If the null 296 hypothesis is rejected at significance level α (where typically α is taken to be 5%), the new 297 null hypothesis becomes a model with two creep regimes present and this is tested against the 298 alternative of a model with three regime changes, using once again a three step procedure. That 299 300 is:

Step 1: Regress y on $\{1, x_1, x_2, w_1, wx_1, w_1x_2\}$ (i.e. assuming one regime change as in Eq, [3d]) to get estimates of the residuals u_2 shown in Eq. [3d].

303 Step 2: Regress u_2 on $\{1, x_1, x_2, w_1, wx_1, w_1x_2, x^2_1, x^3_1, x_1x_2, x_1x^2_1, x_1x^3_1, x^3_1x_2\}$.

Step 3: Compute the Lagrange multiplier statistic $LM = NR^2$ where N is the sample size and R² is the coefficient of determination from the regression carried out in step 2. Under the null hypothesis of one regime change, LM is asymptotically distributed as a chi square variable with 6 degrees of freedom. Accept the model with three different creep regimes if the null hypothesis is rejected at significance level $\tau\alpha$, $0 < \tau < 1$. Reducing the significance level compared to the preceding test favours parsimonious models. Choosing τ is left to the modeller, but $\tau = .5$ is a common choice.

311

This sequential estimation and testing is continued until the first acceptance of the null hypothesis. This yields the specification for the final model and determines the number of creep mechanisms generating the experimental creep data.

316

IV. APPLICATIONS

317 A. 1Cr - 1Mo - 0.25V Steel

In order to apply the sequential testing procedure described in section III above to the 1Cr - 1Mo - 0.25V data shown in Fig. 1a, it is necessary to first construct the residuals u_1 in Eq. [2]. Columns 2 and 3 of Table I show the estimates made for the parameters in Eq. [2]. The t values show that all the parameters are statistically significant at the 1% significance level with the value for d implying an activation energy of just over 300kJmol⁻¹. These estimates imply that the residuals u_1 are given by

324
$$u_1 = y - (23.2390 - 6.9101x_1 - 304.4505x_2)$$
 [6a]

325 The second and third columns of Table II show the results obtained when u_1 is regressed on $\{1, x_1, x_2, x^{2}_1, x^{3}_1, x^{4}_1, x_1x_2, x^{2}_1x_2, x^{3}_1x_2\}$ for the second step of the three step test procedure. 326 It reveals that the parameter in front of x_1x_2 in Eq. [5c] is statistically different from zero at the 327 5% significance level. Eq. [5c] also suggests that this can only be so if there is a statistically 328 significant change in the activation energy (as then $d_2 - d_1 \neq 0$). This part of Table II also reveals 329 330 that the parameter in front of x_1 in Eq. [5c] is statistically different from zero at the 5% significance level. Again, Eq. [5c] also suggests that this can only be so if there is a statistically 331 significant change in either b_1 or a_1 or both, as then $b_2 - b_1 \neq 0$, and or $a_2 - a_1 \neq 0$. This is true 332 because by using the residual u_1 on the left hand side of Eq. [5c] instead of y, the parameter b_1 333 334 is "pulled" (i.e. should be zero in the regression) from this equation during the regression. All these t tests are consistent with the estimates made of the parameters in Eq. [3d] to be discussed 335 further below. 336

As shown in the second and third columns of Table II the R^2 value is quite high at just over 64%. Consequently, the chi square variable (TR²), that test the null hypothesis of no change in creep regime, is statistically significant even at the 1% significance level, meaning that the null hypothesis of just one creep mechanism can be rejected. Thus, there are at least two different creep mechanisms generating the minimum creep rates shown in this 1Cr - 1Mo - 0.25V data set.

To test for the presence of a third creep mechanism, it is necessary to next construct the residuals u₂ in Eq. [3d]. The last two columns of Table I shows the estimates made for the parameters in Eqs. [3b,3d]. These estimates imply that the restricted residuals u₂ are given by

346
$$u_2 = y - (24.4028 - 4.3613x_1 - 317.5693x_2 - 10.3971w - 1.5702wx_1 + 82.0410wx_2)$$
 [6b]

347 with w given by

348
$$w = \frac{1}{1 + \exp[-17.7071(x_1 - 0.4656)]}$$
 [6d]

The last two columns of Table II shows the results obtained when u_2 is regressed on {1, x₁, x₂, x²₁, x³₁, x⁴₁, x₁x₂, x²₁x₂, x³₁x₂} for the second step of the three step test procedure. It reveals that none of the parameter in Eq. [5c] are statistically different from zero (even at the lo% significance level). It is not surprising therefore that the R² value is very low at just over 1% so that the chi square variable (TR²), that test the null hypothesis of one change in creep regime, is statistically insignificant (even at the 10% significance level). Thus the null hypothesis of just one creep regime change cannot be comprehensively rejected. Thus, there are exactly two different creep mechanisms generating the minimum creep rates recorded within this 1Cr - 1Mo - 0.25V data set.

358 As briefly mentioned above, the last two columns of Table II shows the results obtained when the modified Wilshire model applied to the 1Cr - 1Mo - 0.25V data shown in Fig. 1a with 359 two competing creep mechanisms. These estimates are slightly at odds with those originally 360 stated by Wilshire and Scharning and as summarised in Fig. 1a. The last two columns of Table 361 I reveals that the break appears to occur at a normalised stress of 0.47. Whilst this is slightly 362 higher than the value provided by Wilshire and Scharning (0.4), the main differences between 363 364 their results and those shown in this paper stem from the value for $\beta_1 = 17.7$ shown in Table I. This relatively low value gives rise to the sigmoidal curve shown in Fig.2. As can be seen from 365 this figure, a transition from a low to a high stress regime occurs not instantaneously at a 366 normalised stress of 0.47, but very gradually over a wider normalised stress range. At a 367 368 normalised stress of 0.47, w in Eq. [3b] equals 0.5 implying that deformation is equally governed by two competing creep mechanisms. However, once the normalised stress falls to 369 0.3, deformation is predominantly determined by one of these mechanisms (w = 0.05 implies 370 95% determined) and once the normalised stress reaches about 0.6, deformation is 371 predominantly determined by the other mechanism (w = 0.95 implies 95% determined). For 372 373 this modified model to be equivalent to Wilshire's original specification, β_1 would need to be quite large (over 500) so that then the sigmoidal function in Fig.2 would become very step -374 essentially giving a very sharp and rapid transition between these two regimes. 375

The values for d_1 and d_2 shown in the last two columns of Table I help interpret what 376 377 these competing creep mechanisms might be. At normalised stress below 0.3, w is less than 0.05 in value implying that the values for a_1 , b_1 and d_1 in Eq. [3a] are predominant in describing 378 the minimum creep rate. The value for d_1 in particular implies an activation energy of 379 approximately 320kJmol⁻¹. At normalised stress above 0.6, w is more than 0.95 in value 380 381 implying that the values for a_2 , b_2 and d_2 in Eq. [3a] are predominant in describing the minimum creep rate. The value for d_2 in particular implies an activation energy of approximately 382 230kJmol⁻¹. Furthermore, these activation energies are statistically significantly different from 383 each other at the 1% significance level (as shown by the student t values in the d_2 - d_1 row and 384 third column of Table I). This is consistent with the results shown in the first half of Table II 385 386 which showed the parameter in front of x_1x_2 to be statistically significant - when using u_1 are the regressor variable. 387

388 This result is very different from the original Wilshire and Scharning paper where the activation was quoted to be 300kJmol¹ at all levels of the normalised stress. This varying 389 activation energy must also cast doubt on their explanation for the kink in the best fit line shown 390 391 in Fig.1a. For creep to occur predominantly by diffusion controlled generation and movement of dislocations within the lattice structure only, (with particle coarsening within the lattice 392 393 being the cause of changing k_2 and v values), no matter what the stress level is, the activation energy should also be unchanging with respect to stress. Neither can the changing values for 394 395 k_2 and v be attributable to a change from creep occurring from the generation of new 396 dislocations within the lattice structure itself to creep occurring from the movement of 397 dislocations pre-existing in the grain boundary zones only. Because then the activation energy 398 would be lower at low normalised stress. This is not in agreement with the estimates made from the data where the opposite appears to be true - the sigmoidal curve shown in Fig.3a shows the 399 activation energy increasing with decreasing normalised stresses. However, the activation 400 energies shown in Fig. 3a are consistent with the traditional view that Nabarro –Herring 401 402 diffusional creep becomes more dominant at lower stresses. This is further supported by the fact that in the NIMS data set the lower stress tests are at the highest temperatures. If this is so, 403 then 320kJmol⁻¹ would be that activation energy for self-diffusion. The only way to explain the 404 lower activation energy that is estimated for the high stress regime (which in the NIMS data 405 set also corresponds to low temperatures), is to suggest that under this condition the dominant 406 creep mechanism is preferential diffusion along dislocations (without dislocation movement) 407 or coble creep, i.e. stress directed vacancy flow along grain boundaries . 408

409 There is also a statistically significant difference between b_1 and b_2 and between a_1 and a_2 as revealed by the student t values in Table I (in the a_2 - a_1 and b_2 - b_1 rows). Thus the gradual 410 411 switch in the deformation mechanism with stress is also associated with changing values for both k_2 and v. In Figs. 3a,b the values for Q^*_{c} , k_2 and v are multiplied by the changing value 412 for w shown in Fig.2 to give an impression of how these parameters change with the normalised 413 stress. As can be seen, the main changes in the values for these parameters takes place over the 414 normalised stress range of 0.3 to 0.6. It is over this stress range then that the deformation 415 416 mechanism driving creep switches. These changes drive the shape of the solid curve in Fig. 3a. Along the stretch a – b we have the familiar negative relationship between $\ln(-\ln(\sigma/\sigma_T s))$ and 417 $\ln[\dot{\epsilon}_m \cdot \exp{\{Q^*_c/RT\}}]$. Then Q^*_c starts to change rapidly and this leads to the stretch of the curve 418 between b and c. Finally, over the normalised stress range 0.1 - 0.3, the familiar negative 419 420 relationship between $\ln(-\ln(\sigma/\sigma_T s))$ and $\ln[\dot{\epsilon}_m \cdot exp\{Q^*c/RT\}]$ returns but now the activation energy is much higher than before. 421

422 B. 2.25Cr-1Mo steel

In order to apply the sequential testing procedure described in section III above to the 2.25Cr-1Mo data shown in Fig. 1b, it is necessary to first construct the residuals u_1 in Eq. [2]. Columns two and three of Table III shows the estimates made for the parameters in Eq. [2]. The t values show that parameters b and d are statistically significant at the 1% significance level with the value for d implying an activation energy of nearly 200kJmol⁻¹. These estimates imply that the residuals u_1 are given by

$$u_1 = y - (7.4906 - 6.1446x_1 - 190.2616x_2)$$
[7a]

Columns 2 and 3 of Table IV show the results obtained when u_1 is then regressed on {1, x_1 , x_2 , x^{2_1} , x^{3_1} , x^{4_1} , x_1x_2 , $x^{2_1}x_2$, $x^{3_1}x_2$ } for the second step of the three step test procedure. It reveals that at the 5% significance level, the only parameter to be statistically insignificant is that in front of x^{4_1} in Eq. [5c]. Eq. [5c] also suggests that this result can only be so if there is a statistically insignificant change in the value for v in Eq. [2] (as then $b_2 - b_1 = 0$). Eq. [5c] also suggests that the statistical significance of all the other parameters shown in these two columns can only be so if there is a statistically significant change in both d_1 and a_1 . Then $a_2 - a_1 \neq 0$ 437 (leading to the parameter in front of x_1, x^2_1 and x^3_1 in the first half of Table IV being significantly 438 different from zero), and $d_2 - d_1 \neq 0$ (leading to the parameter in front of x_1x_2 in the first half of 439 Table IV being significantly different from zero). All these t tests are consistent with the 440 estimates made of the parameters in Eq. [3d] to be discussed further below.

As shown in columns two and three of Table IV, the R^2 value is quite high at just over 91%, so that the chi square variable (TR²), that test the null hypothesis of no change in creep regime, is statistically significant even at the 1% significance level. This in turn means that the null hypothesis of just one creep mechanism is rejected by the data. Thus, there are at least two different creep mechanisms generating the minimum creep rates recorded in this 2.25Cr-1Mo data set.

447 To test for the presence of a third creep mechanism, it is necessary to next construct the 448 residuals u_2 in Eq. [3d]. The middle section of Table III shows the estimates made for the 449 parameters in Eq. [3d]. These estimates imply that the residuals u_2 are given by

450
$$u_2 = y - (6.5490 - 4.8317x_1 - 198.9690x_2 + 29.4426w + 0.0727wx_1 - 165.6330wx_2)$$
 [7b]

451 with w given by

452

$$w = \frac{1}{1 + \exp[-38.6887(x_1 - 0.2638)]}$$
[7c]

The last two columns of Table IV show the results obtained when u_2 is regressed on {1, 453 $x_1, x_2, x_1^2, x_1^3, x_1^4, x_1x_2, x_1^2x_2, x_1^3x_2$ for the second step of the three step test procedure. It 454 reveals, first of all, that the parameter in front of x^{4}_{1} is statistically different from zero at the 455 5% significance level. Eq. [5c] also suggests that this can only be so if there is a statistically 456 significant change in the slope of the best fit line in Fig. 1b (as then $b_2 - b_1 \neq 0$). The statistical 457 significance of the parameter in front of x^{2}_{1} at the 5% significance level may also indicate that 458 the intercept of the best fit line in Fig. 1b changes (as then $a_2 - a_1 \neq 0$). The last two columns 459 Table IV also reveals the parameter in front of $x_{1}^{2}x_{2}$ is statistically different from zero at the 460 10% significance level. Eq. [5c] also suggests that this can only be so if there is a statistically 461 significant change in the activation energy (as then $d_2 - d_1 \neq 0$). These t statistics are therefore 462 suggestive that for this material at least three creep mechanisms are at work. This is further 463 confirmed by the R² value, which is quite high at just over 34% so that the chi square variable 464 (TR²), that test the null hypothesis of just two creep regimes, is statistically significant at the 465 10% significance level. This in turn means that the null hypothesis of just two creep mechanism 466 is rejected by the data. Thus, there are at least three different creep mechanisms generating the 467 468 minimum creep rates recorded in this 2.25Cr - 1Mo steel data set.

Although the results of testing the null hypothesis of exactly three creep regimes using this LM test are not shown here, the test leads to the acceptance of this null hypothesis - even at the 10% significance level. Thus for this material there appears to be three distinctly different creep regimes or mechanisms and the parameter estimates of Eq. [4b,c] shown in the last two columns of Table III throw some light on the nature of these regimes. These estimates are slightly at odds with those originally stated by Wilshire and Whittaker - which are shown in Fig. 1b. The last two columns of Table III reveals that the two break points appear to occur at normalised stresses of 0.26 and 0.42. Whilst these are slightly different to the values provided by Wilshire and Whittaker (around 0.1 and 0.5 respectively), the main difference stems from the values for $\beta_1 = 50.5$ and $\beta_3 = 18.1$ shown towards the bottom of Table III. These values give rise to the sigmoidal and bell shaped curves shown in Fig.4.

480 As can be seen from this figure, a transition from a low to a medium stress regime and then from a medium to a high stress regime occurs, but not instantaneously, at normalised 481 stresses of 0.26 and 0,42 respectively. Below a normalised stress of 0.3, about 90% of the 482 deformation is governed by the first creep mechanism (summarised by the value for w_1). The 483 remaining deformation is governed by the other two mechanisms. Then at a normalised stress 484 of around 0.30 the second mechanism dominates with about 80% of the deformation being 485 controlled by this mechanism (as shown by the value for w_2). Beyond a normalised stress of 486 0.42, the third mechanism starts to dominate with around 90% of the deformation being 487 governed by this last mechanism at normalised stresses of 0.55 and above (as reflected in the 488 value for w₃). For this modified model to be equivalent to Wilshire and Whittaker's original 489 specification, β_1 and β_3 would need to be quite large (over 500) so that then the sigmoid al 490 functions in Fig.4 would become very step, and the bell shaped function very compressed,-491 essentially giving a very sharp and rapid transition between the regimes. 492

The values for d_1 , d_2 and d_3 in the last two columns of Table III help interpret what 493 these competing creep regimes or mechanisms might be. At a normalised stress around 0.30, 494 w_2 is about 0.8 in value implying that the values for a_2 , b_2 and d_2 in Eq. [4b] are predominant 495 in describing deformation and the minimum creep rate. The value for d_2 then implies an 496 497 activation energy of approximately 240kJmol¹ which is consistent with the estimates made by Wilshire and Whittaker for this middle stress regime (see Fig. 1b where the activation energy 498 is given by the authors at 230kJmol¹). At normalised stresses less than 0.2, w₁ is 0.8 or more 499 in value implying that the values for a_1 , b_1 and d_1 in Eq. [4b] are predominant in describing 500 501 deformation and the minimum creep rate. The value for d_1 shown in the last two columns of Table III then implies an activation energy of approximately 200kJmol⁻¹, but because the t 502 statistic on d_1 - d_2 is insignificant (implying d_1 - d_2 is insignificantly different from zero), the 503 conclusion must be that the activation energy in this low stress regime is not different to that 504 in the medium stress regime. This is very different to the conclusion given by Wilshire and 505 Whittaker who maintain that the activation energy is much higher in this low stress regime (but 506 they provide no statistical proof for this hypothesis). At normalised stresses above 0.55, w₃ is 507 0.8 or more in value implying that the values for a_3 , b_3 and d_3 in Eq. [4b] are predominant in 508 describing deformation and the minimum creep rate. The value for d₃ shown in Table III then 509 implies an activation energy of approximately 400kJmol^{-1} , and because the t statistic on d₃-d₂ 510 is statistically significant, the conclusion must be that the activation energy in this high stress 511 regime is different to that in both the medium and low stress regimes. This activation energy is 512 much higher than that quoted by Wilshire and Whittaker who maintain that the activation 513 514 energy is around 280kJmol⁻¹ in this high stress regime (see Fig. 1b).

515 According to Wilshire and Whittaker ^[5], when $\sigma > \sigma_Y$, creep is controlled by the 516 generation and movement of dislocations within the grains. This would require a high 517 activation energy, which is consistent with the result described above where an activation energy is estimated at around 400kJmol⁻¹. In contrast, when $\sigma < \sigma_Y$, Wilshire and Whittaker 518 suggest that dislocations are not generated within the grains. Instead, creep occurs within the 519 520 grain boundary zones, i.e. by grain boundary sliding and or diffusion along existing 521 dislocations and grain boundaries. This requires a lower activation energy, which is consistent with the result described above where an activation energy of around 230kJmol¹ is estimated 522 523 for medium stresses. Wilshire and Whittaker then suggest another change in creep and creep rupture behaviour occurs when σ approximately equals $0.2\sigma_{TS}$. With this material, they suggest 524 the original ferrite/bainite microstructure degrades to ferrite and molybdenum carbide particles 525 in long term tests at the highest creep temperatures, with very coarse carbide particles forming 526 along the grain boundaries (which takes place in long-term tests at the highest creep 527 temperatures). This then enables deformation to once again be determined by processes within 528 the lattice structure, where the activation energy is greatest. Whilst the results in this paper 529 suggest that a mechanism change does indeed occur in the transition from medium to very low 530 stresses, there is no significant increase in the activation energy. Contrary to the Wilshire 531 532 explanation, this result suggests that creep is not predominantly determined by processes occurring within the lattice structure material - because the activation energy is highest within 533 the bulk. It would seem instead that sliding and or diffusion along existing dislocations and 534 grain boundaries still predominates at these very low stresses. But that the coarsening of the 535 536 carbide particles reduces creep strength further given the different stress relation shown in the low stress regime compared to the medium stress regime, i.e. allows creep rates to be much 537 higher than would be predicted using relations that apply in the medium stress regime. 538

There are also statistically significant differences between b₁, b₂ and b₃ and between a₁, 539 a_2 and a_3 as revealed by the student t values in Table III (in the a_1 - a_2 , a_3 - a_2 and the b_1 - b_2 , b_3 - b_2 540 rows). Thus the gradual switch in deformation mechanisms with stress is also associated with 541 changing values for both k_2 and v. In Figs. 5a,b the values for Q^*_{c} , k_2 and v are multiplied by 542 the changing value for w_i shown in Fig. 4 to give an impression of how these parameters change 543 with the normalised stress. As can be seen, the main changes in the values for these parameters 544 takes place over the normalised stress range of 0.2 to 0.5. k₂ appears to continually increase 545 546 with the normalised stress, whilst v is similar in value at the highest and lowest stresses with a temporary increase over the intermediate normalised stress ranges. The values for v at the end 547 points (i.e. at points a and d in Fig. 5a) are very similar to the estimates made by Wilshire and 548 Whittaker in their original study – as can be seen by a comparison of Fig.1a with Fig.5a. 549 However, the values for k_2 in this study appear a little larger in comparison. 550

Finally, the solid curves in Fig.6 shows what the predictions given in Fig.5 look like in 551 stress - minimum creep rate space. It can be seen that the predictions trace out well defined 552 smooth curves as the stress level varies. In contrast to this, the dotted "curves" show the 553 predictions obtained when the weighting functions w_1 to w_3 are step like in nature which then 554 closely corresponds to the original Wilshire - Whittaker specification for this material. The 555 predictions at some of the temperatures are very discontinuous due to abrupt changes in the 556 activation energy and the functional relationship of the minimum creep rate with stress. These 557 discontinuities do not make physical sense and lead to rather bizarre behaviour. For example, 558

at 873K (600°C) and between 53 MPa and 41 MPa, the minimum creep rate slows down in a uniform fashion, but then just before a stress of 41 MPa is reached the model predicts the creep rate will suddenly increase even though there has been little change in the stress level. From a creep perspective this makes little sense and reflects the incorrect specification of the way the activation energy changes with stress (in reality it is gradual rather than abrupt).

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V. CONCLUSIONS

This paper has put forward a statistical test for determining the correct number of discontinuities to use within the Wilshire equations and also a method for allowing these discontinuities to change more gradually with the normalised stress level - so that the methodology is more in line with the accepted view as to how creep mechanism evolve with changing test conditions. The new findings obtained using this modified methodology include:

In their study of 1Cr - 1Mo - 0.25V steel, Wilshire and Scharning worked with a constant 570 i. activation energy of 300kJmol⁻¹ and a change in the relationship between the minimum 571 572 creep rate and the normalised stress that occurred abruptly at a normalised stress of 0.4. In contrast, this paper found that the activation energy also changed with the normalised 573 stress. Further, these changes occurred gradually over a normalised stress range of around 574 0.3 to 0.6. This changing activation energy in turn casts doubt on the authors view that 575 the changing values for k_2 and v were the result of particle coarsening associated with 576 long test durations at lower stresses. 577

In their study of 2.25Cr-1Mo Wilshire and Whittaker worked with an activation energy 578 ii. that was lower for mid-range normalised stresses (230kJmol⁻¹) than it was for any other 579 580 value of the normalised stress (where they took the activation energy to be 280 kJmol^{-1}). 581 In contrast, this paper found the activation energy to be around 400kJmol¹ at the highest values for the normalised stress but around 240kJmol⁻¹ for all other values of the 582 normalised stress. These difference suggest that creep is not predominantly determined 583 by processes occurring within the lattice structure at these lowest stress values as 584 585 originally suggested by these authors. Over the normalised stress range 0.2 to 0.5, creep is predominantly determined by a single process with an activation energy of 240kJmol⁻ 586 ¹. Below a normalised stress of 0.25, there is quite an abrupt change in the values for k_2 587 and v, whilst in contrast, the changes in k₂ and v are more gradual for increases in the 588 589 normalised stress above a value of 0.45.

When the new procedures outlined in this paper were applied to 2.25Cr-1Mo steel, they
 produce more accurate and realistic looking long term predictions of the minimum creep
 rate.

593 An important area for future work includes applying the methodology outlined in this 594 paper to other steel alloys to confirm whether this approach also produced better long term 595 predictions for these materials, and better understanding of the changing deformation 596 mechanisms.

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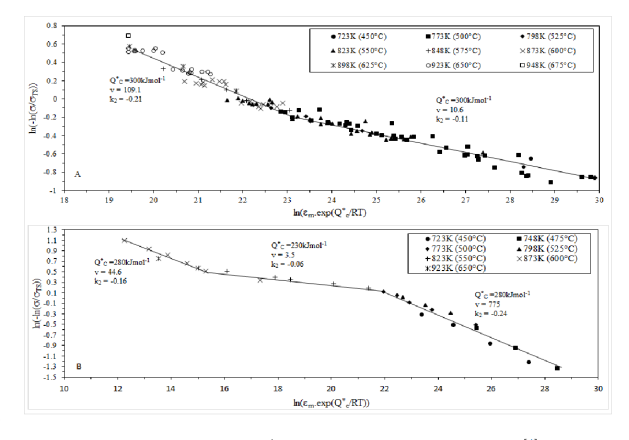


Fig.1 – The best values for k_2 , v and Q^*_c determined by Wilshire and Scharning^[4] and Wilshire and Whittaker^[5] were found by plotting $\ln[\epsilon_m.exp(Q^*_c/RT)]$ against $\ln(-\ln(\sigma/\sigma_{TS}))$ for a. 1Cr – 1Mo – 0.25 steel forgings for rotors and shafts and b. for 2.25Cr-1Mo steel tubes.

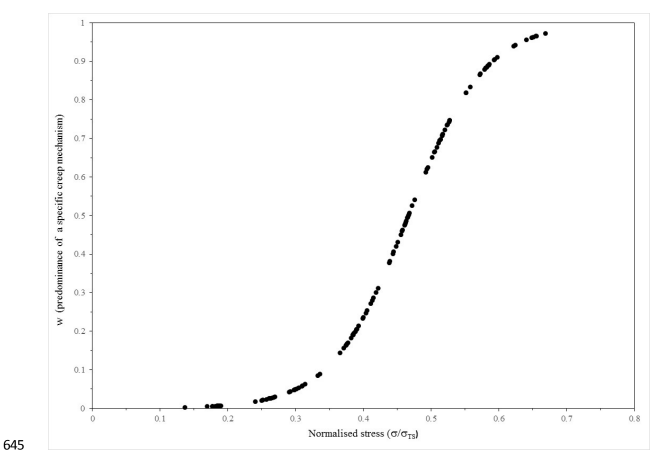


Fig. 2 - The dominance of two different deformation mechanisms at different stresses for 1Cr1Mo-0.25V steel.

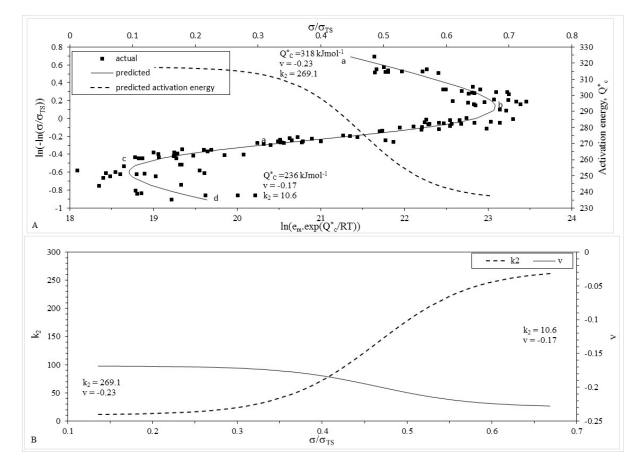


Fig.3 – Dependence of a. $\ln[\epsilon_m.exp(Q^*c/RT)]$ on $\ln(-\ln(\sigma/\sigma_{TS}))$ and the activation energy on the normalised stress and b. dependence of k_2 and v on the normalised stress for 1Cr-1-Mo-0.25V steel at various temperatures.

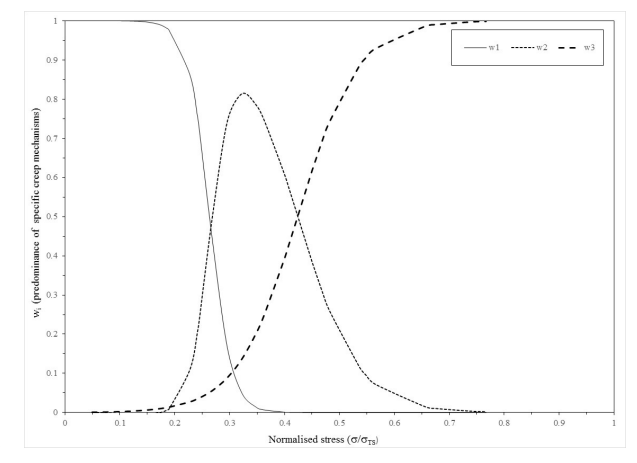


Fig. 4 - The dominance of three different deformation mechanisms at different stresses for2.25Cr-1Mo steel.

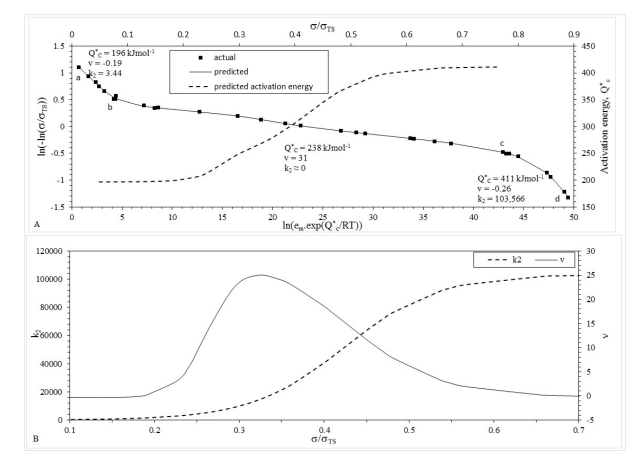


Fig.5 – Dependence of a. $\ln[\epsilon_m.exp(Q^*c/RT)]$ on $\ln(-\ln(\sigma/\sigma_{TS}))$ and the activation energy on the normalised stress and b. dependence of k_2 and v on the normalised stress for 2.25Cr-1Mo steel at various temperatures.

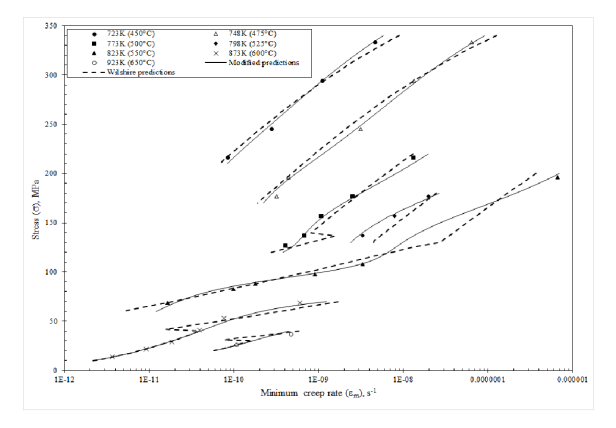




Fig.6 – Minimum creep rates in 2.25Cr-1Mo steel tubes predicted by the modified and original
 specifications of the Wilshire equation.

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	Eq.	[2]	Eqs. [3b,3d]		
Parameters	Least squares estimates	t value	Least squares estimates	t value	
a	23.2390	14.43***	-	-	
b	-6.9101	-24.24***	_	-	
d	-304.4505	-26.89***	-	-	
a1	-	-	24.4028	13.22***	
b 1	-	-	-4.3613	-10.09***	
d_1	-	-	-317.5693	-24.44***	
a ₂ -a ₁	-	-	-10.3971	-2.91***	
b ₂ -b ₁	-	-	-1.5702	-2.67***	
d ₂ -d ₁	-	-	82.0411	3.34***	
a2	-	-	14.0057	6.06***	
b ₂	-	-	-5.9316	-11.35***	
d ₂	-	-	-235.5282	-8.47***	
x [*] 1	-	-	0.4656	-	
β1	-	-	17.7071	-	

Table I. Least Squares Estimates for the Parameters in Eq. [2] and Eqs. [3b,3d] when using 1Cr-1Mo-0.25V Steel Forging Data

***Parameters are statistically different from zero at the $\alpha = 1\%$ and above significance level. ** Parameters are statistically different from zero at the $\alpha = 5\%$ and above significance level. * Parameters are statistically different from zero at the $\alpha = 10\%$ and above significance level. T is the sample size (T=121). t has a student t distribution with T - 3 degrees of freedom for Eq. [2] and T - 6 degrees of

freedom for Eq. [3d].

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	u1		u ₂		
Variable	Least squares estimates	t & Chi square values	Least squares estimates	t & Chi square values	
Constant	-0.9344	-0.54	-0.7656	-0.45	
X1	14.8038	2.51**	0.8635	0.15	
X2	2.9373	0.24	5.3381	0.44	
x ² 1	6.3926	0.38	17.1725	1.01	
x ³ 1	-0.3812	-0.02	17.3221	0.82	
x ⁴ 1	-4.1614	-1.17	-2.7544	-0.78	
X1X2	-98.4011	2.31**	-5.6813	-0.13	
$x^2 x_1 x_2$	-25.0805	-0.21	-122.0521	-1.03	
x ³ 1x ₂	-7.2728	-0.04	-139.0271	-0.85	
R ² (%)	64.17	-	1.03	-	
$LM = TR^2$	-	77.68***	-	1.25	

Table II. Results from Regressing u_1 and u_2 on $\{1, x_1, x_2, x^{2}_1, x^{3}_1, x^{4}_1, x_1x_2, x^{2}_1x_2, x^{3}_1x_2\}$ 681 when using 1Cr-1Mo-0.25V Steel Forging Data

***Parameters are statistically different from zero at the $\alpha = 1\%$ and above significance level. **Parameters are statistically different from zero at the $\alpha = 5\%$ and above significance level. *Parameters are statistically different from zero at the $\alpha = 10\%$ and above significance level.

 R^2 is the coefficient of determination or the percentage variation in u explained by all the variables shown in the first column of the table.

 TR^2 has a chi square distribution with 6 degrees of freedom. T is the sample size (T = 121). t has a student t distribution with T - 9 degrees of freedom.

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	Eq.	[2]	Eq. [3d]		Eq. [4b,c]	
Parameters	Estimate	t value	Estimate	t value	Estimate	t value
a	7.4906	1.15	-	-	-	-
b	-6.1446	-7.99***	-	-	-	-
d	-190.2616	-4.38***	-	-	-	_
a2	-	-	-	-	15.6319	3.16***
b ₂	-	-	-	-	0.0325	0.02
d ₂	-	-	-	-	-238.3443	-7.45***
a ₁ -a ₂	-	-	-29.4426	-7.58***	-8.9704	-1.65*
a3-a2	-	-	-	-	28.8024	4.43***
b1-b2	-	-	-0.0727	0.14	-5.422	-3.81***
b ₃ -b ₂	-	-	-	-	-3.8803	-2.98***
d ₁ -d ₂	-	-	165.6330	6.33***	42.1756	1.18
d3-d2	-	-	-	-	-172.4897	-4.18***
a ₁	-	-	6.5419	2.06^{**}	6.6615	19.24***
a ₃	-	-	-	-	44.3426	3.18***
b1	-	-	-4.8317	-9.59***	-5.3897	-12.28***
b ₃	-	-	-	-	-3.8478	-15.54***
d1	-	-	-198.9690	-9.05***	-196.1688	-6.58***
d3	-	-	-	-	-410.8314	-28.42***
x*1	-	-	0.2638	-	0.2638	-
β1	-	-	38.6887	-	50.5172	-
X ^{**} 1	-	-	-	-	0.4239	-
β ₃	-	-	-	-	18.1292	-
** Parameters	are statistically are statistically re statistically e size $(T = 31)$ t distribution	different f different fi different fro).	rom zero at the	$\alpha = 5\%$ and $\alpha = 10\%$ and	l above signifie l above signifie	cance leve cance leve

Table III. Least Squares Estimates for the Parameters in Eq. [2] and Eqs. [3b,3d] whenusing 2.25Cr - 1Mo Steel Tube Data

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Table IV. Results from Regressing	u_1 and u_2 on $\{1, x_1,$	$x_2, x_1^2, x_1^3, x_1^4, x_1x_2, x_1^2x_2, x_1^3x_2$
when using	2.25Cr - 1Mo Steel	Tube Data

	uı		u ₂	
Variable	Least squares estimates	t & Chi square values	Least squares estimates	t& Chi square values
Constant	10.8126	2.80**	3.1162	-1.35
X1	-50.3627	-6.05***	-7.7382	-1.55
X2	-62.4319	-2.47**	20.0285	1.32
x ² 1	22.5831	12.09***	14.9435	2.07**
x ³ 1	47.5139	3.63***	5.7176	0.72
x ⁴ 1	-0.9135	-0.84	-1.3998	-2.13**
X1X2	315.8915	5.60***	51.5805	1.53
$x^{2}x^{2}x^{2}$	-162.0437	-2.09**	-88.0569	-1.90*
x ³ 1X2	-314.1496	-3.45***	-44.4357	-0.83
R ² (%)	91.03	-	34.25	-
TR ²	-	28.22***	-	10.72*

***Parameters are statistically different from zero at the $\alpha = 1\%$ and above significance level. **Parameters are statistically different from zero at the $\alpha = 5\%$ and above significance level. *Parameters are statistically different from zero at the $\alpha = 10\%$ and above significance level.

 R^2 is the coefficient of determination or the percentage variation in u explained by all the variables shown in the first column of the table.

 TR^2 has a chi square distribution with 6 degrees of freedom. T is the sample size (T = 31). t has a student t distribution with T - 9 degrees of freedom.

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