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Multi-objective parameter estimation for simulating canopy transpiration in forested watersheds

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Abstract

A Jarvis based [Philos. Trans. R. Soc. London, Ser. B 273 (1976) 593] model of canopy stomatal conductance was evaluated in context of its application to simulating transpiration in a conifer forest covered watershed in the Central Sierra Nevada of California, USA. Parameters influencing stomatal conductance were assigned values using Monte Carlo sampling. Model calibration was conducted by evaluating predicted latent heat fluxes against thermal remote sensing estimates of surface temperature. A fuzzy logic approach was used to select or reject simulations and form a restricted set of ensemble parameter solutions. Parameter estimates derived from the ensembles were evaluated using theory on how stomatal conductance regulates leaf water potential to prevent runaway cavitation. Canopy level parameters were found to be sufficient for predicting hydraulically consistent transpiration when soils were well watered. A rooting length parameter controlling the amount of plant available water was a sufficient addition to the parameter set to predict hydraulically consistent transpiration when soil moisture stress was occurring. Variations in maximum stomatal conductance among different hillslopes within the watershed were explained by a light threshold parameter. The results demonstrate that the Jarvis model can be reliably parameterized using thermal remote sensing data for estimating transpiration in meso-scale watersheds.

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Keywords: Stomatal conductance; Simulation model; Parameter estimation; Transpiration; Thermal remote sensing

1. Introduction

Spatially variable transpiration is a major component flux simulated by distributed land surface process models. Although there is a scarcity of observational data to directly support large-scale simulation of canopy transpiration from forests, many models operating from watersheds to global scales (Running and Coughlan, 1988; Aber and Federer, 1992; Band et al., 1993; Running and Hunt, 1993; Famiglietti and Wood, 1994; Wigmosta et al., 1994; Vertessy et al., 1996; Foley et al., 1996, 2000; Mackay and Band, 1997; Sellers et al., 1997; and others) simulate transpiration using some form of the Penman-Monteith combination equation (P-M) (Monteith, 1965) and one of several empirical models

HYDROL 6850-30/4/2003-12:33-MUKUND-69041 - MODEL 3

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97 of stomatal conductance (Jarvis, 1976; Lohammar et al., 1980; Ball et al., 1987). Although there are 98 numerous sources of uncertainty in P-M models, the 99 unknowns associated with stomatal conductance are 100 most critical for understanding vegetation responses 101 to climate change and land use pressures. Mackay 102 et al. (2003) show that Jarvis-type models mimic the 103 stomatal regulation of leaf water potentials in a 104 detailed canopy model operating at sub-daily time 105 resolution, with in situ micrometeorological measure-106 107 ments and sap flux data for calibration. The study by Mackay et al. (2003) was limited to four forest stands 108 each approximately 30 m in diameter. To extend the 109 applicability of their approach to watershed or global 110 scales necessitates finding tractable data sources and 111 more moderate model detail. 112

One potential source of data for characterizing the 113 state of a land surface is thermal remote sensing. 114 Thermal remote sensing data is a measure of surface 115 temperature, which is a function of energy partition-116 ing and surface resistance. Foliage temperature has 117 been shown to relate to soil moisture, plant moisture 118 stress, and transpiration (Idso et al., 1978; Jackson 119 et al., 1981). This has led to numerous applications of 120 121 thermal remote sensing to study canopy processes (Goward et al., 1985; Pierce and Congalton, 1988; 122 Holbo and Luvall, 1989; Nemani and Running, 1989; 123 Kustas et al., 1994; Carlson et al., 1995; Anderson 124 et al., 1997; Franks and Beven, 1997; Bastiaanssen 125 et al., 1998; Norman et al., 2000). These methods have 126 been successful using very different approaches. 127 Representative approaches include modeling with 128 detailed ground flux measurements and sub-daily 129 remote sensing measurements to look at thermal 130 inertia (Anderson et al., 1997; Norman et al., 2000), 131 132 limiting the analysis to large-scale remote sensing with strong moisture gradients and well-coupled 133 forest canopies (Nemani and Running, 1989), or 134 calibrating a model using Monte Carlo analysis with 135 contrasting vegetation types (Franks and Beven, 136 1997). 137

In this paper, the latter two approaches are
combined to evaluate a Jarvis-type model as used
within the Regional Hydro-Ecological Simulation
System (RHESSys) (Band et al., 1993) applied to a
meso-scale watershed in the Central Sierra Nevada.
RHESSys has been and continues to be applied
successfully in numerous forested watershed studies

(Band et al., 1996; Creed et al., 1996; Watson et al., 145 1996; Mackay and Band, 1997; White et al., 1998; 146 Baron et al., 2000; Tague and Band, 2001; Zhu and 147 Mackay, 2001; Mackay, 2003; Samanta and Mackay, 148 2003). Its ability to predict stream flow, nitrate export, 149 and responses to forest manipulation and road 150 construction are documented in these studies. How-151 ever, neither RHESSys nor any model like it has been 152 tested for consistent spatial transpiration. The ques-153 tion is whether RHESSys can produce transpiration at 154 a watershed scale that is consistent with plant water 155 relation theory. Significant progress has recently been 156 made in the understanding of hydraulic limitations to 157 canopy transpiration. These limitations have helped in 158 constraining parameters for a Jarvis-type model at the 159 stand level (Mackay et al., 2003). It is hypothesized 160 that transpiration estimates can be similarly evaluated 161 through calibration of Jarvis-type parameters control-162 ling stomatal conductance at larger spatial scales and 163 on daily time-step, by comparing model predictions 164 and thermal remote sensing estimates of surface 165 energy partitioning. Rejecting this hypothesis would 166 mean that the model is either deficient with respect to 167 processes such as soil water distribution, the Jarvis-168 type model is a poor approximation to the stomatal 169 conductance at daily time-steps or at larger spatial 170 scales, or the combination of data and models used in 171 the analysis are insufficient to resolve the underlying 172 regulation of leaf water potential by stomata, which 173 ultimately affects energy partitioning. 174

The paper is organized as follows. First, a summary 175 of the most relevant elements of plant water relations 176 are presented to build a framework for relating the 177 Jarvis-type stomatal conductance model to tree 178 hydraulic theory. This is followed by the methods, 179 which describes an automated calibration based on 180 Samanta and Mackay (2003) and restricted using the 181 tree hydraulic theory (Mackay et al., 2003). Relevant 182 details on RHESSys are presented, followed by the 183 analysis of thermal remote sensing imagery, results, 184 discussion, and conclusions. 185

2. Plant water relations and the Jarvis model of stomatal conductance

Canopy transpiration from forests is generally 191 simulated with the well-known P–M 'big leaf' model 192

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(Monteith, 1965). A key variable in P-M is canopy 193 conductance, which is determined by the product of 194 stomatal conductance, $g_{\rm S}$ and leaf area, L. Models of 195 stomatal conductance address one or more environ-196 mental (extrinsic) and physiological (intrinsic) con-197 ditions of the leaf stomata. There are two distinct 198 empirical models of stomatal conductance: Ball-199 Berry (Ball et al., 1987; Leuning et al., 1995) and 200 Jarvis (Jarvis, 1976; Lohammar et al., 1980). The 201 Ball-Berry model emphasizes the rate of carbon 202 assimilation in controlling stomatal conductance, but 203 places little emphasis on water supply and demand. 204 Jarvis models more directly address water supply and 205 demand through soil water limitation and atmospheric 206 vapor pressure deficit functions, respectively. Jarvis-207 type models have the general form: 208

$$\begin{array}{ll} 210 & g_{\rm S} = g_{\rm Smax} \prod f_i, \\ 211 & \end{array} \tag{1}$$

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where g_{Smax} is a theoretical maximum stomatal 212 conductance under assumed optimal environment 213 and leaf conditions. A series of multiplicative 214 functions of environmental factors (f_i) are applied to 215 reduce actual leaf level stomatal conductance from the 216 theoretical maximum level. Typically, one function 217 218 considers the stomatal sensitivity, δ , to atmospheric vapor pressure deficit, D. The g_{Smax} parameter can 219 vary widely among and within species (Kelliher et al., 220 1995; Ewers et al., 2001) and δ is widely believed to 221 222 increase with maximum stomatal conductance (Jarvis, 1976, 1980; McNaughton and Jarvis, 1991; Monteith, 223 1995; Saliendra et al., 1995; Oren et al., 1999). Low 224 225 vapor pressure gradient conditions favor stomatal control by assimilation rate, but as D increases 226 stomata close to reduce water loss (Ball et al., 1987; 227 Monteith, 1995; Saliendra et al., 1995; Yong et al., 228 229 1997). One advantage of the Jarvis model for water flux estimates is that it directly addresses plant 230 response to D. This suggests that it operates best 231 when the rate of water loss is high and, hence, 232 hydrologically significant. Furthermore, recent devel-233 opments in plant hydraulic theory have been success-234 fully combined with Jarvis models (Oren et al., 1999; 235 Ewers et al., 2000) and tested in diurnal canopy 236 models on a range of forest species (Mackay et al., 237 2003). 238

Eq. (1) is a proxy for the hydraulic functioning of the soil-plant-atmosphere continuum. In the absence of stomatal control, a high rate of water loss from a 241 plant leads to a rapid decline in leaf water potential. 242 This increases the risk of hydraulic failure in the plant 243 (Sperry et al., 1998; Oren et al., 1999; Ewers et al., 244 2000). The relationship between stomatal conduc-245 tance and water potential can be expressed with the 246 steady-state assumption and Darcy's law (Tyree and 247 Ewers, 1991; Ewers et al., 2000): 248

$$G_{\rm S} = K_{\rm L} / D(\Psi_{\rm S} - \Psi_{\rm L} - h\rho_{\rm w}g),$$
 (2) 250

251 where G_S is canopy average leaf level stomatal 252 conductance, $K_{\rm L}$ is leaf-specific whole-plant 253 hydraulic conductance, $\Psi_{\rm S}$ and $\Psi_{\rm L}$ are bulk soil 254 and leaf water potentials, respectively, and $h\rho_w g$ is 255 the gravitational potential for a plant of height h. 256 $K_{\rm L}$ declines with water potential due to soil drying, 257 cavitation in the xylem, and other factors. As $K_{\rm L}$ 258 declines a further drop in water potential is needed 259 to sustain increasing $E_{\rm C}$ per unit leaf area (L). This 260 drop in water potential further reduces $K_{\rm L}$ as a 261 positive feedback. Once the maximum safe tran-262 spiration rate is exceeded then runaway cavitation 263 ensues. Runaway cavitation refers to $K_{\rm L}$ being 264 driven to zero causing hydraulic failure and 265 possible plant mortality (Tyree and Sperry, 1988). 266 Eq. (2) also captures the well-known fact that G_S is 267 inversely proportional to D (Jarvis, 1976; 268 McNaughton and Jarvis, 1991), which drives 269 down stomatal conductance even when plants are 270 well watered in the soil. This is shown concep-271 tually in Fig. 1. $G_{\rm S}$ is further shown to be sensitive 272 to increasing D in proportion to some maximum 273 conductance, g_{Smax} , or its proxy (Oren et al., 1999): 274

$$G_{\rm S} = G_{\rm Sref} - m \ln D, \qquad (3) \qquad (3)$$

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where G_{Sref} is a substitute for g_{Smax} defined at D =279 1 kPa and $m = dG_S/d \ln D$ is the sensitivity of 280 stomatal conductance to increasing D. In Eq. (3), 281 G_{Sref} is typically near linearly related to the 282 sensitivity of stomatal conductance to D. Based 283 on a large amount of porometry and sap flux data, 284 Oren et al. (1999) have shown that $m \approx 0.6G_{\text{Sref}}$ 285 applies universally to all species whose stomata 286 regulate leaf water potential to just prevent run-287 away cavitation. 288

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Fig. 1. The rates of stomatal closure (lines) in response to vapor pressure deficit are proportional to the maximum stomatal conductances (circles) (a), have been shown to result in all species that just regulate leaf water potential to prevent runaway cavitation to lie along a line representing the stomatal sensitivity to vapor pressure deficit versus reference conductance at 1 kPa (ln D = 0).

306 3. Methods

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308 3.1. Overview of the model parameterization309 approach

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In this paper, two distinct stages of parameter 311 estimation are used. The first stage of parameter 312 estimation follows a traditional automated calibration. 313 Many simulations are run in which parameters 314 affecting stomatal conductance are assigned values 315 using Monte Carlo sampling. Each simulation result is 316 then evaluated by applying a linear least-squares 317 318 regression between simulated evaporative fraction from RHESSys and surface temperature from thermal 319 remote sensing data. For each least-squares 320 regression, the coefficient of determination (or R^2) is 321 calculated. The R^2 is then considered a fuzzy measure 322 of the goodness-of-fit for its respective simulation 323 result. The set of R^2 measures for all simulations is 324 considered a fuzzy set (Samanta and Mackay, 2003). 325 An information-theoretic tool is then applied to the 326 fuzzy set to form a restricted set in which only 'good' 327 simulations retained. A restricted set is used as an 328 ensemble solution in the second stage of parameter 329 estimation. A separate ensemble solution is produced 330 for each areal unit simulated by RHESSys, and in the 331 present work these areal units are hillslopes. 332

The second stage of parameter estimation applies the universal line represented by Eq. (3) and shown in Fig. 1. For each ensemble δ is related to g_{Smax} . When a δ/g_{Smax} combination falls on the universal line, it is assumed to be consistent with plant hydraulic theory. Otherwise, it is considered inconsistent with the theory. To be consistent with the conventions of Oren et al. (1999), Jarvis parameters (δ and g_{Smax}) are mapped into their respective universal line counterparts (*m* and G_{Sref}). For the remainder of this paper the use of δ and g_{Smax} will be restricted to discussion of model function, not plant physiology. Discussion of plant physiology will use *m* and G_{Sref} .

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3.2. Stage 1: model-independent automated parameterization

An objective automated parameter estimation 367 framework (Samanta and Mackay, 2003) was used 368 to calibrate RHESSys. The approach is based on a 369 number of hydrologic parameter estimation schemes 370 (Kuczera, 1982, 1983; Sorooshian and Gupta, 1983; 371 Spear and Hornberger, 1990; van Stratten and Kees-372 man, 1991; Klepper et al., 1991; Binley and Beven, 373 1991; Melching, 1995; Kuczera and Parent, 1998; 374 Gupta et al., 1998; Yapo et al., 1998; Boyle et al., 375 2000). It combines Monte Carlo sampling and 376 measures of uncertainty derived from information 377 processing. One information-theoretic expression of 378 uncertainty in information processing is the Hartley 379 (1928) Function: 380

$$H(A) = \log_2|A|, \tag{4} \quad \begin{array}{c} 381\\ 382\\ 382 \end{array}$$

where H(A) is the Hartley Function for a finite set, A, 383 and |A| is its cardinality. Eq. (4) is a measure of 384

the non-specificity arising from an inability to identify a unique solution. Higher values of H(A) represent greater non-specificity. If A represents a set of retained simulations, each defined by a simulation model and its parameters, then H(A) is the nonspecificity associated with this equifinal (Beven and Binley, 1992) set of simulations.

The above notion of the non-specificity of a set of 392 simulations can be extended to incorporate measures 393 of model fitness. If each simulation has an associated 394 measure of fitness or degree of fit to some observation 395 data, then the acceptable set of models can be 396 considered a fuzzy set (Zadeh, 1965). For a fuzzy 397 set, F, defined by a measurement of fitness, $f(x_i)$, for 398 member x_i within the domain, X, of all simulations, a 399 fuzzy logic measure of the non-specificity of F is 400 (Higashi and Klir, 1982; Klir and Wierman, 1998): 401

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$$U(F) = \int_{0}^{h(F)} \log_2 |\alpha F| d\alpha + (1 - h(F)) \log_2 |X|,$$
 (5)

405 where U(F) is the U-uncertainty associated with F, 406 $|^{\alpha}F|$ is the cardinality of an α -cut of F (i.e. the number 407 of members that remain in the set if all members with 408 a degree of fitness less than α are removed from F), 409 h(F) is the height of F (or maximum degree of fitness 410 in F), and |X| is the cardinality of the universal set (i.e. 411 the population of simulations). A discrete approxi-412 mation of Eq. (5) is given by 413

$$U(r) = \sum_{i=2}^{n} (r_i - r_{i+1}) \log_2 i + (1 - r_1) \log_2 n,$$

$$(6)$$

where r is the ordered possibility distribution (Zadeh, 417 1978) derived from the fuzzy set F and r_{n+1} is 418 assumed to be 0. Fig. 2 shows two typical possibility 419 distributions relating $f(x_i)$, which is the goodness-of-420 fit measure, and $|^{\alpha}F|$, as well as the physical meaning 421 of the α -cut. Both distributions are shown to have an 422 α -cut of about 0.6, at which they yield very different 423 cardinalities. Relations that are skewed towards the 424 low end, and thus have only a few high $f(x_i)$ models, 425 are better than relations having too many high $f(x_i)$ 426 values. The ideal distribution of simulations relation 427 428 has one with $f(x_i) = 1.0$ and all others with $f(x_i) =$ 0.0. This would represent a case where the best 429 simulation is uniquely identifiable. For numerous 430 reasons, including model flaws, parameter trade-offs, 431 432 and limitations of objective functions, hydrologic



Fig. 2. Shown is an illustration of the types of possibility 450 distributions that are commonly obtained from the U-uncertainty 451 analysis. The plots represent the relationship between cardinality on 452 the y-axis and value returned by some model evaluation objective function applied to simulations of two hypothetical landscape 453 patches. Shown are the α -cuts obtained by minimizing |U(r) - U(r)| = 0454 $\log_2 k$. It is important to note that similar α -cuts may be obtained 455 from different sets of simulations in which the retained model sets 456 have very different cardinalities. 457

models do not satisfy this ideal (Oreskes et al., 1994; Mackay and Robinson, 2000).

It is tempting to set the α -cut at such a level that 460 only one simulation is retained. This is the traditional 461 approach to model calibration. However, an objective 462 criterion provides a better way to define the α -cut to 463 form a restricted set from the fuzzy set (Samanta and 464 Mackay, 2003). Initially, this α -cut should be selected 465 with caution. On the one hand, useful information in 466 the fuzzy set may be lost if an arbitrarily high α -cut is 467 selected. An arbitrarily high α -cut may also admit a 468 false sense of specificity to the identified model 469 parameters. On the other hand, an arbitrarily low α -470 cut may include too many simulations of poor quality. 471 Information Theory presents a rich set of tools for 472 extracting the full information content of a fuzzy set. 473 One tool, the Principle of Uncertainty Invariance (Klir 474 and Wierman, 1998), transforms a fuzzy set into a 475 'crisp' restricted set that approximates the respective 476 fuzzy set by virtue of having the same U-uncertainty. 477 The α -cut is chosen at the *k*th element of the ordered 478 possibility distribution at which $|U(r) - \log_2 k|$ is 479 minimized. As an illustration, consider the ordered 480

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fuzzy set $F = \{0.9, 0.8, 0.8, 0.7, 0.6, 0.4, 0.1, 0.1\}$. It is 481 possible to calculate $|^{\alpha}F| = 4.4$ as the sum of fuzzy 482 memberships in the set and U(r) = 2.2 as its 483 associated U-uncertainty. The size of the restricted 484 set is determined by equating the U-uncertainty of the 485 fuzzy set (Eq. (6)) to the Hartley Function (Eq. (4)) for 486 the desired crisp set. The value of k (5 in this example) 487 is the required cardinality of the restricted set, which 488 means that the top five elements in F are retained and 489 the α -cut is placed at 0.6. An advantage of this 490 approach is that the selection of the α -cut is not 491 subject to interpretation or modification as the goals of 492 a modeling exercise change. A disadvantage of the 493 approach is that it does not consider intuition about 494 the physical system, which is an essential part of 495 parameter estimation (Boyle et al., 2000). However, 496 once an objective solution set has been established, 497 further analysis can be applied to determine if a more 498 refined solution can be found. In this paper, a simple 499 averaging of the respective parameter values from the 500 restricted sets is tested by evaluating the degree to 501 which the chosen parameters for a Jarvis-type model 502 are consistent with the plant hydraulic theory 503 relationships (Refer to Section 2). The Jarvis-type 504 model and its parameterization are described in 505 506 Section 3.3.

3.3. Stage 2: parameterization specific to RHESSyscanopy transpiration

RHESSys combines forest canopy gas exchange 511 processes, soil moisture balance, and lateral saturated 512 through flow (Band et al., 1993). It represents 513 watersheds as collections of hillslopes, which are 514 themselves divided into elevation zones for adiabatic 515 adjustment of air temperature, T_a . Each elevation zone 516 517 is segmented into hydrologically uniform patches 518 defined using the TOPMODEL topography and soil index (TSI) (Beven and Kirkby, 1979; Beven, 1986; 519 Sivapalan et al., 1987; Quinn et al., 1995). All vertical 520 fluxes, including evapotranspiration, are calculated at 521 the patch level. Complete details on the design and 522 implementation of RHESSys are provided in previous 523 publications (Band et al., 1993; Mackay and Band, 524 1997; Mackay, 2003). This section focuses on model 525 components that directly affect the calculation of 526 stomatal conductance. Leaf level stomatal conduc-527 tance $(m s^{-1})$ is determined (from Eq. (1)) in 528

RHESSys as: 529

 $g_{\rm S} = g_{\rm Smax} g_1(D) g_2(\Psi_{\rm L}) g_3(Q), \tag{7} \qquad \begin{array}{c} 530\\ 531 \end{array}$

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where

 $g_1(D) = 1 - \delta D$ (8) $533 \\ 534$

and $\delta [(kPa)^{-1}]$ has been previously defined as the 535 sensitivity of stomatal conductance to D. From the 536 earlier discussion it should be apparent that Eq. (8) is a 537 surrogate for stomatal response to the rate of water 538 loss from the canopy, which is strongly related to D in 539 well-ventilated forest canopies (McNaughton and 540 Jarvis, 1991). This can be interpreted in the terms 541 used in Eq. (4) by mapping g_{Smax} and δ into G_{Sref} and 542 $dG_S/d \ln D$, respectively. The g_{Smax} parameter rep-543 resents a theoretical optimal stomatal conductance 544 under ideal conditions (i.e. low D, sufficient light, 545 moderate temperature, well-watered soils). Following 546 Eq. (8) $g_{\rm S} \equiv g_{\rm Smax}$ at D = 0 kPa. It is not possible to 547 measure $g_{\rm S}$ at D = 0 kPa, and so $g_{\rm Smax}$ is transformed 548 into G_{Sref} (at D = 1 kPa as defined by Oren et al. 549 (1999)) as follows: 550

 $G_{\text{Sref}} = g_{\text{Smax}}(1 - \delta).$ (9) 551 552

Similarly δ is converted into *m* as follows:

 $m = \mathrm{d}G_{\mathrm{S}}/\mathrm{d}\ln D = g_{\mathrm{Smax}}\mathrm{d}\delta/\mathrm{d}\ln D, \qquad (10) \qquad \begin{array}{c} 554\\ 555 \end{array}$

in which the derivative is calculated by finite-556 difference at two values of D (Mackay et al., 2003). 557 The other functions $(g_2 \text{ and } g_3)$ in Eq. (7) are not 558 modified from their standard expressions. Stomatal 559 conductance is further reduced using a hyperbolic 560 function of leaf water potential, $\Psi_{\rm L}$, which is assumed 561 to be equal to pre-dawn soil water potential calculated 562 using a van Ganuchten (1980) formulation parameter-563 ized using Brooks and Corey (1964) soil hydraulic 564 parameters. Stomatal conductance is reduced linearly 565 with the ratio of absorbed radiation in the canopy to a 566 minimum threshold radiation, Q_{\min} . 567

RHESSys calculates soil water potential within a 568 depth of soil defined by plant rooting length, $R_{\rm L}$. 569 $R_{\rm L}$ is varied spatially to account for two competing 570 controls on water supply: (1) the tendency for 571 capillarity to recharge a drying rooting zone when 572 there is a shallow perched water table, and (2) the 573 need for sufficient $R_{\rm L}$ to support the water demand 574 associated with a given leaf area index, L (Grier 575 and Running, 1977; Gholz, 1982), or, equivalently, 576

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the root-shoot ratio. $R_{\rm L}$ is spatially co-varied with L 577 and S (Mackay, 2003), expressed for a given patch, 578 *i*, of uniform TSI within hillslope, *h*, as 579

$$\sum_{582}^{581} R_{\mathrm{L},i} = R'_h L_{h,i} f_h(S_{h,i}), \qquad (11)$$

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584 where R'_h is intrinsic rooting length (per unit $L_{h,i}$) 585 for a mesic site on h (m), $f_h(S_{h,i})$ is a function, 586 which, through dimensional analysis, describes 587 relative soil saturation deficit at *i* with respect to 588 average soil saturation deficit within h, and $L_{h,i}$ is 589 leaf area index defined at i on h (m² m⁻²). Eq. (11) 590 accounts for higher water demand environments 591 (e.g. south-facing slopes, high temperatures) by 592 increasing R'_h . Furthermore, high $L_{h,i}$ requires 593 proportionally deeper roots to supply adequate 594 water to the canopy. Sites with low saturation 595 deficits do not require deep roots, and roots cannot 596 survive long in a shallow water table (Larcher, 597 1995). 598

The derivation of $f_h(S_{h,i})$ is based on elements of 599 TOPMODEL (Beven, 1986; Beven and Kirkby, 1979; 600 Sivapalan et al., 1987). Profile saturation deficit is 601 scaled from hillslope facet average saturation deficit 602 as 603

607 where $\langle S_h \rangle$ is mean saturation deficit for h (m), $S_{h,i}$ is 608 saturation deficit of patch iof h(m), m_h is a parameter 609 that describes the rate of decay of saturated hydraulic 610 conductivity through a soil profile in hillslope h, TSI_{*h*,*i*} 611 is topography-soils index at patch i of hillslope h (m), 612 and $\langle TSI_h \rangle$ is hillslope average topography-soils 613 index (m). $TSI_{h,i}$ is calculated as follows:

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₆₁₇ TSI_{*h,i*} = ln
$$\left(\frac{a_{h,i}t_h}{t_{h,i}\tan\beta_{h,i}}\right)$$
, (13)

where $a_{h,i}$ is accumulated upslope drainage area per 620 unit contour width (m), $\beta_{h,i}$ is local topographic slope, 621 $t_{h,i}$ is local soil transmissivity (m day⁻¹), and t_h is 622 hillslope average transmissivity (m day $^{-1}$). Eq. (12) 623 can be conditionally expanded to describe $S_{h,i}$ as 624

function of
$$\langle S_h \rangle$$
:

$$\int = \langle S_h \rangle, \quad \frac{\langle \mathrm{TSI}_h \rangle}{\mathrm{TSI}_{h,i}} = 1$$

$$S_{h,i} = \begin{cases} \langle \langle S_h \rangle, & \frac{\langle \mathrm{TSI}_h \rangle}{\mathrm{TSI}_{h,i}} < 1. \end{cases}$$
(14)
$$\begin{array}{c} 629\\ 630\\ 631 \end{array}$$

From Eq. (14) the relative saturation deficit can be determined from hillslope average conditions with the dimensionless form:

$$f_h(S_{h,i}) = \frac{\langle \mathrm{TSI}_h \rangle}{\mathrm{TSI}_{h,i}}.$$
(15)
$$\begin{array}{c} 638\\ 639\\ 640 \end{array}$$

641 Substituting Eq. (15) into Eq. (11) gives a relation for 642 scaling rooting length along joint moisture and leaf 643 area gradients within a hillslope, 644

$$R_{L_{h,i}} = R'_h L_{T_{h,i}} \frac{\langle \text{TSI}_h \rangle}{\text{TSI}_{h,i}}.$$
(16)

The parameter, R'_h , can be modified to linearly scale the volume of soil available for plant access to water.

3.4. Evaluation data set

Data to test the Jarvis-based model in RHESSys 653 was obtained from the Onion Creek Experimental 654 Forest, a 10 km² watershed located along the crest of 655 the Central Sierra Nevada of Annual precipitation 656 averages 1300 mm, of which 90% falls as snow 657 between October 1 and March 31. Relief within the 658 watershed is about 1000 m, with the highest elevation 659 occurring at ca. 2600 m above mean sea level. Soils 660 are generally poorly developed loamy sands overlying 661 highly permeable rhyolitic ash and latite deposits 662 (MacDonald, 1987). The soils and underlying parent 663 material have a high storage capacity producing long 664 lags in the baseflow recession (MacDonald, 1986). 665 Vegetation cover is predominantly mature to old-666 growth mixed needle-leaf conifers. The dominant tree 667 species in the basin are White Fir (Abies concolor), 668 Red Fir (Abies magnifica), Sugar Pine (Pinus 669 lambertiana), Jeffrey Pine (Pinus jeffreyi), Lodgepole 670 Pine (Pinus contorta), and Incense Cedar (Calocedrus 671 decurrens). Cedar generally occupies lower 672

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elevations, pine at mid-elevations, and fir at higher elevations.

Radiometric thermal data was acquired with the 675 NASA-Ames Research Center's airborne Thematic 676 Mapper Simulator (TMS), a Daedelus scanner flown 677 aboard a U-2 aircraft on July 2 and August 6, 1985. 678 The image data was collected in conjunction with 679 field-measured surface temperature and soil moisture 680 (Pierce and Congalton, 1988; MacDonald, 1989). On 681 July 2, soil moisture measurements in the watershed 682 showed soil moisture levels between field capacity 683 and saturation. On August 6 the soils were well below 684 field capacity and approaching wilting point in some 685 plots (MacDonald, 1989). The TMS images were 686 collected at approximately solar noon at a flying 687 height commensurate with a sensor resolution of 30 m 688 (or 0.09 ha per observation). The flight lines were 689 centered over Onion Creek to avoid off-nadir 690 geometry problems, allowing for relatively accurate 691 co-registration of the TMS with a Landsat Thematic 692 Mapper 4 (TM) scene (RMS error <15 m). Sensor 693 radiance values, r, were converted to kinetic tem-694 peratures, t, with the following regression and 695 parameters (Table 1) obtained from black-body 696 radiation sensor calibration reference plates, atmos-697 pheric correction, and ground control within Onion 698 Creek (Pierce and Congalton, 1988): 699

$$\begin{array}{l} 700\\ 701\\ 702 \end{array} \quad T_{\rm s} = \frac{t_{\rm max} - t_{\rm min}}{r_{\rm max} - r_{\rm min}} (10r - r_{\rm min}) + t_{\rm min}. \end{array}$$
(17)

No adjustment was made for variations in surface emissivity, as the range of cover types in Onion Creek (occasional bare soil to predominantly dense conifer canopy) should have emissivities of 0.9-0.95, which translates into 1-3% error when inverting the Stefan– Boltzmann Equation to convert from radiant to kinetic temperature.

Kinetic temperatures are influenced by a number
of other factors in complex topography. These
factors include stomatal, boundary and aerodynamic
Table 1
Black body reference plate regression parameters used in Eq. (14) to
convert TMS pixel values to radiant temperatures

Date	t _{max}	t _{min}	r _{max}	$r_{\rm min}$
July 2	32.21	8.18	149.00	105.00
August 6	32.32	8.49	150.00	110.00

resistances, which effect latent and sensible heat 721 transfer; meteorological differences, such as cloud 722 cover and air temperature between flight dates; 723 temperature lapse with elevation; land surface-sun 724 geometry; and forest canopy density. Meteorological 725 data was obtained from the Central Sierra Snow Lab. 726 Fig. 3 shows daily temperatures (maximum and 727 728 minimum) and precipitation for the period beginning before the July 2 TMS scene and ending after the 729 730 August 6 scene. Both dates show similar daily high 731 and low temperatures, and clear skies following 732 several days with no precipitation. Aerodynamic 733 conductance was assumed high (set at 0.2 m s^{-1}) for 734 the dense, conifer needle-leaf forests covering much of Onion Creek. TM data was used to derive leaf area index (LAI) using ground-based calibration and image processing procedures described in Nemani



Fig. 3. This shows the meteorological conditions spanning the period around the two selected dates (July 2 and August 6) for the TMS imagery. Note that both dates occur during a dry period with no recent rainfall, and with similar maximum and minimum temperatures.

et al. (1993). A USGS 30 m digital elevation model 769 (DEM) was used to identify hillslopes that account for 770 variability in land surface-sun geometry, and 771 elevation bands that account for temperature lapse 772 773 rates. A digitized soils map was acquired from the US 774 Forest Service, and used for soil parameterization of 775 RHESSys. Table 2 summarizes topography, veg-776 etation, and other hillslope facet mean properties of 777 Onion Creek, partitioned into 25 hillslope partitions 778 (Liang and Mackay, 2000). 779

A separate objective function was used for each 780 TMS scene for each model. Linear least-squares 781 regression between evaporative fraction, $\sigma = \lambda E_{\rm C}/Q$, 782 and surface departure from air temperature, dT =783 $T_{\rm s} - T_{\rm a}$ was used to evaluate model fitness. The 784 reason for using σ is that it remains nearly constant 785 during the mid-day period (Crago, 1996; Crago and 786 Brutsaert, 1996) and so it is less likely to be sensitive 787

to short-term changes in micrometeorological con-817 ditions than is latent heat flux alone. Furthermore, 818 whereas $\lambda E_{\rm C}$ and Q both give a non-linear fit to 819 surface temperature data due to their asymptotic 820 response with leaf area index through the Beer-821 Lambert Equation, their ratio is close to linear. As a 822 requirement for using TMS imagery collected at a 823 single point in time at solar noon, it is necessary to 824 825 assume that midday dT is correlated with daily $E_{\rm C}$ at a given aerodynamic conductance (Seguin and Itier, 826 827 1983; Carlson et al., 1995). Furthermore, it was 828 assumed that soil heat flux had a minimal influence on 829 canopy temperature given the high leaf area (Carlson 830 et al., 1995). Finally, residuals derived from the best 831 model for each hillslope facet for each TMS scene 832 were checked for trends. There were no visible trends 833 in the residuals that would suggest a linear fit was 834 inappropriate. 835

Table 2

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790 This shows a summary of the average properties of all hillslopes in the Onion Creek watershed partitioning. Number of zones refers to the 791 number of elevation zones used to capture within-hillslope adiabatic temperature lapse rates. Each zone represents an increment of 30 m 792 altitude, such that a hillslope with 20 zones has an elevation range of about 600 m

Hillslope identifier	Area (ha)	Aspect (°)	Elevation (m)	Slope (°)	$L ({\rm m}^2{\rm m}^{-2})$	$\langle \mathrm{TSI}_h \rangle$ (m)	Number of zones
1	30.6	1/3 1	1701 3	11.3	8 50	8.2	11
2	34.4	265.1	1804 3	10.2	9.02	7.26	12
3	64.7	111.1	1896.4	12.8	9.17	8.11	10
4	21.0	264.4	1834.1	9.3	9.13	6.67	4
5	11.6	98.3	1917.0	13.6	8.98	7.69	9
6	1.9	159.2	1864.9	15.0	8.92	8.63	3
7	30.1	91.6	2041.3	10.2	9.11	7.47	11
8	24.5	187.5	2063.9	11.2	10.57	6.83	12
9	82.7	133.3	2108.7	15.5	8.53	7.12	15
10	43.7	219.8	2074.8	13.7	8.43	7.08	15
12	2.7	293.3	1851.1	9.6	7.89	6.87	3
13	159.2	119.6	2060.7	16.8	7.46	7.13	18
14	92.2	238.6	2091.8	18.6	8.77	6.99	19
15	7.9	206.9	1880.2	3.2	7.75	10.6	3
16	67.9	283.7	1958.0	12.7	8.65	7.59	12
17	42.7	140.0	2130.7	19.3	6.50	6.57	17
18	54.1	229.3	2144.2	18.3	7.29	6.71	18
19	6.8	182.6	1972.8	11.9	5.82	7.93	4
20	34.8	284.4	2052.2	15.0	8.30	7.2	11
21	76.7	200.3	2229.5	19.4	6.23	6.6	17
22	21.4	274.4	2257.7	21.5	4.20	6.46	20
23	43.6	237.2	2185.4	18.5	6.21	6.72	20
24	57.4	300.7	2214.6	20.7	5.31	6.23	18
25	123.1	203.9	1993.3	9.9	9.23	7.64	21
26	195.5	276.5	2112.2	12.9	8.46	7.64	22

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865 **4. Results**

866 Three sets of simulations were run with different 867 parameter combinations. Following Mackay et al. 868 (2003), an initial set of 2000 simulations were run 869 for each of the 25 hillslopes with values for three 870 canopy conductance parameters, g_{Smax} (range 17-871 $126 \text{ mmol m}^{-2} \text{ s}^{-1}$), Q_{\min} (range 3000 -872 9000 kJ m⁻² day⁻¹), and δ (range 0.07–0.74 kPa⁻¹) 873 randomly sampled from a naive (uniform) distri-874 bution. For this first set of simulations no parameters 875 controlling available soil water were adjusted. Table 3 876 summarizes the average of these parameters taken 877 from the ensemble of models retained using |U(r)| – 878 $\log_2 k$ for the respective July and August TMS scenes. 879 The Q_{\min} averages are generally higher for August 880 than for July, but no such clear pattern exists for g_{Smax} 881 or δ . This indicates that there is a need for greater 882 883 reduction in stomatal conductance for the August date in comparison to July, and this reduction is not 913 satisfied with a reduction in either g_{Smax} or δ . This is 914 further evident in Fig. 4, which shows how stomatal 915 sensitivity to D versus reference conductance for each 916 hillslope compares to the theoretical universal line 917 (Refer to Fig. 1) of stomatal regulation of leaf water 918 potential (Oren et al., 1999). The figure clearly shows 919 that the simulations for July are closely following the 920 universal line. However, the simulations for August 921 show considerable scatter, indicating that the simu-922 lation model is performing poorly when judged 923 against the plant water-relations theory. 924

One hypothesis for the poor model performance for August is that soil moisture limitation is not adequately simulated. This reasoning would make sense if hillslopes at low elevation, with south-facing aspects and high leaf area index tended to follow the universal line the least. Fig. 5 shows a relationship between the calculated slope (m/G_{Sref}) and leaf area

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Summary of ensemble averages for the respective parameters for calibration with the July and August TMS images, respectively

Hillslope identifier	July			August		
	$\frac{E(Q_{\min})}{(\text{kJ m}^{-2} \text{ day}^{-1})}$	$\frac{E(G_{\rm Smax})}{(\rm mmol \ m^{-2} \ s^{-1})}$	$\frac{E(\delta)}{((kPa)^{-1})}$	$\frac{E(Q_{\min})}{(\text{kJ m}^{-2} \text{ day}^{-1})}$	$\frac{E(G_{\rm Smax})}{(\rm mmol \ m^{-2} \ s^{-1})}$	$E(\delta)$ $((kPa)^{-1})$
1	6565.0	85.8	0.433	5449.6	99.1	0.540
2	5641.0	73.8	0.410	5827.2	72.0	0.401
3	6305.2	80.0	0.411	7182.0	74.4	0.466
4	5639.8	75.9	0.420	6269.5	71.9	0.489
5	5892.9	73.2	0.389	6086.8	72.4	0.465
5	4890.0	43.9	0.359	5055.3	89.3	0.411
7	5823.0	68.6	0.428	7221.1	75.7	0.452
8	3588.6	61.0	0.406	5630.6	82.0	0.409
9	6689.6	67.4	0.433	6376.3	88.0	0.421
10	5798.1	71.3	0.403	6866.2	102.3	0.384
12	6655.6	61.4	0.442	3809.2	81.5	0.305
13	3561.4	41.9	0.413	7721.5	80.0	0.454
14	4910.7	54.8	0.442	7631.9	81.0	0.516
15	4112.3	60.7	0.411	5207.2	65.2	0.348
16	6006.6	80.8	0.402	6891.3	81.0	0.447
17	5503.4	69.1	0.412	5748.1	64.2	0.412
18	5558.6	71.4	0.417	5288.4	65.2	0.397
19	4738.1	44.6	0.413	4923.1	76.7	0.349
20	3460.6	66.8	0.406	3798.1	75.8	0.391
21	5353.8	81.2	0.432	5094.9	71.3	0.439
22	5475.0	84.6	0.396	5508.0	84.6	0.410
23	5451.5	77.1	0.394	5233.5	66.5	0.394
24	5371.6	81.9	0.416	5537.8	83.8	0.402
25	4010.8	43.5	0.420	6989.5	71.7	0.497
26	3917.9	59.3	0.415	6842.6	75.0	0.449

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Fig. 4. These plots show the relationships between stomatal sensitivity to vapor pressure deficit and reference conductance at a vapor pressure deficit of 1 kPa for all hillslopes for both July and August TMS scene dates. The dashed line has a slope $0.6 \ln (kPa)^{-1}$, which has been shown by a large volume of data and modeling to represent a universal tradeoff in stomatal function (Oren et al., 1999; Ewers et al., 2000). Plants that regulate their water potential to just prevent runaway cavitation should theoretically fall on this line. The solid lines represent linear regressions for the July and August points, respectively. For the August plots the regression line was forced through the origin, since we expect stomatal sensitivity to be zero at full stomatal closure. Note that both fitted lines have slope near the theoretical 0.6 line, and in particular the July result shows a strong fit ($R^2 = 0.89$).

index, L, for the hillslopes organized into three groups 988 by elevation. At low elevation, high evaporative 989 demand, sites there is a strong positive relationship 990 between the m/G_{Sref} and L. This relationship is weaker 991 at the intermediate elevations and non-existent at the 992 high elevations. Where m/G_{Sref} is near 0.6 indicates 993 that the simulations are consistent with the universal 994 line. When m/G_{Sref} exceeds 0.6 this indicates δ is 995 increasing to compensate for an underestimated soil 996 997 moisture stress or too low Q_{\min} . At values below 0.6 it is possible that simulated soil moisture stress is too 998 great. While this result supports the notion that soil 999 moisture limits are poorly represented in the simu-1000 lations, it does not fully explain why Q_{\min} increases 1001 for August. Fig. 6 shows a positive relationship 1002 between m/G_{Sref} and Q_{\min} . In addition, the position of 1003 1004 each hillslope (above or below $m/G_{\text{Sref}} = 0.6$) can be explained in terms of properties that affect water 1005 supply versus demand. Lower elevation hillslopes and 1006 1007 a steep, southwest aspect hillslope have $m/G_{\text{Sref}} \gg 0.6$ 1008 and correspondingly higher Q_{\min} . This indicates that

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Fig. 5. Shown in this figure are relations between $m/G_{\rm Sref}$, which1051theory suggests should have a value of 0.6, and leaf area index (L).1052The hillslopes have been sorted by their average elevation into low,
medium and high categories to show that at lower elevations where
temperatures are higher there is a relatively strong correspondence
between the $m/G_{\rm Sref}$ and L. This relationship is not significant at the
high elevations at which most of the hillslopes follow the theory.1051

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Fig. 6. Shown in this figure are relations between m/G_{Sref} , which 1072 theory suggests should have a value of 0.6, and Q_{\min} . It shows a 1073 clear positive relationship that suggests that Q_{\min} is compensating 1074 for factors controlling stomata that are not represented properly in the model, such as pre-dawn soil water potential. Individual 1075 hillslopes and groups of hillslopes, which do not follow water 1076 regulation theory ($m/G_{\text{Sref}} = 0.6$) are distinguished in terms of low 1077 elevation or high radiation load (steep, southwest aspect) for greater 1078 than expected apparent stomatal regulation of water potential, and 1079 low moisture demand versus supply $(L/\langle TSI \rangle \ll 1)$ or northwest aspect for lower than expected apparent stomatal regulation of water 1080 potential. 1081

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areas of higher moisture demand for the amount of 1083 supply are simulated as too wet, which forces the 1084 canopy parameters to compensate by exceeding the 1085 universal line. Hillslopes with low L per unit $\langle TSI_h \rangle$ 1086 (an index of water demand versus supply) or north-1087 west aspect have $m/G_{\text{Sref}} \ll 0.6$ and lower Q_{\min} 1088 values. These hillslopes are simulated as having too 1089 little soil moisture, which requires the canopy 1090 parameters to compensate by falling below the 1091 universal line. 1092

It appears that Q_{\min} is being used incorrectly to 1093 compensate for a poor representation of soil water 1094 limitation on stomatal conductance. Since it is a 1095 surrogate for the light limitation on stomatal opening, 1096 which is species dependent, there is no clear 1097 justification for changing Q_{\min} between July and 1098 August. To remedy this a second set of 2000 1099 simulations per hillslope was run using a single Q_{\min} 1100 of 5237 kJ m⁻² day⁻¹ determined as the average July 1101 Q_{\min} for all hillslopes, and the rooting length 1102 parameter, R'_h , was randomly sampled from an 1103 uninformed distribution with a range of 0.01–0.1 m. 1104

The results are shown in Fig. 7. Overall, there is a 1105 substantial improvement in the fit of each hillslope to 1106 the universal line for both dates. The August date still 1107 shows more scatter. Based on the inset plot in Fig. 7 1108 there is a strong, positive relationship between m/G_{Sref} 1109 and the rooting length. This relationship is similar to 1110 the one shown in Fig. 6, indicating a tradeoff is 1111 occurring between R'_h and Q_{\min} . To test this 1112 hypothesis, a third and final set of 2000 simulations 1113 was run using the ensemble average July Q_{\min} values 1114 from Table 3, for the respective hillslopes. The results 1115 are shown in Fig. 8. The results show a tight clustering 1116 of all hillslopes on the universal line, but at a narrow 1117 range of G_{Sref} . Furthermore, R'_h , shown inset in Fig. 8, 1118 now has a narrow range of values, which means all 1119 hillslopes are converging to a point in parameter 1120 space. 1121

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5. Discussion

The parameter tradeoffs (g_{Smax} versus δ) seen in the 1126 daily simulations of transpiration suggest that the 1127 Jarvis model captures stomatal regulation of water 1128 potential. The tradeoffs seen with this relatively 1129 coarse simulation with sparse data are consistent 1130 with those proposed in theory (Monteith, 1995), 1131 extensively tested against a large amount of porome-1132 try and sap flux data (Oren et al., 1999), and obtained 1133 using a half-hourly model driven by in situ micro-1134 meteorology and calibrated with sap flux data 1135 (Mackay et al., 2003). It is important to realize that, 1136 without this direct comparison to water-relations 1137 theory, it is possible that numerous factors would 1138 have contributed to this apparent physically consistent 1139 behavior of the model. For example, there are 1140 abundant degrees of freedom given the uncertainties 1141 in both model inputs and in deriving surface 1142 temperature from TMS. Furthermore, one should be 1143 cautious in interpreting simulation model parameters 1144 obtained from ensemble averages. The values for the 1145 respective parameters may simply approach the mean 1146 of the a priori distributions from which they are 1147 sampled. This problem can be avoided by applying an 1148 iterative refinement of the parameter values, as was 1149 done in this paper. The fact that a large range of G_{Sref} 1150 is obtained for the first and second set of simulations 1151 (Figs. 4 and 7) is proof that parameter values for each 1152

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1174 deficit of 1 kPa for all hillslopes for both July and August TMS scene dates. This result is from the second set of simulations in which the rooting 1175 1223 length scalar, R', is varied. The dashed line has a slope 0.6 ln (kPa)⁻¹, which has been shown by a large volume of data and modeling to 1224 1176 represent a universal tradeoff in stomatal function (Oren et al., 1999; Ewers et al., 2000). Plants that regulate their water potential to just prevent runaway cavitation should theoretically fall on this line. The solid lines represent linear regressions for the July and August points, respectively. 1177 1225 For the August plots the regression line was forced through the origin, since we expect stomatal sensitivity to be zero at full stomatal closure. 1178 1226 Note that both fitted lines have slope near the theoretical 0.6 line, and in particular the July result shows a strong fit ($R^2 = 0.95$). August has a 1179 1227 good fit for all except two hillslopes. Inset plots show relationships between m/G_{Sref} and the rooting length scalar. 1180 1228

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hillslope are not simply taking on the distribution
means for the respective parameters. Instead, the
results show that by following water-relations theory a
progressive refinement of the Jarvis-type model
parameters is obtained.

The progressive refinement of model parameter 1187 values demonstrates an important distinction between 1188 automated versus knowledge-based parameterization 1189 schemes. Automated schemes (Binley and Beven, 1190 1991; Gupta et al., 1998; Samanta and Mackay, 2003) 1191 tend to be naïve and extremely sensitive to the type of 1192 objective functions used, as would be the case in the 1193 present paper. Knowledge-based schemes are sensi-1194 tive to the application domain (Franks and Beven, 1195 1997; Boyle et al., 2000; Mackay et al., 2003), take 1196 into consideration intuition about the function of the 1197 modeled system, and tend to be less sensitive to 1198 1199 specific objective functions. The universal line (Oren 1200 et al., 1999) summarizes a large amount of data on

1229 plant hydraulic function. Based on the results in this 1230 paper for watershed-scale modeling and those of 1231 Mackay et al. (2003) for stand-level modeling, there is 1232 sufficient justification for argument that the Jarvis 1233 model parameters can be made to follow the universal 1234 line. This does not confirm that the models mimic 1235 plant hydraulic function, anymore than it confirms a 1236 direct correspondence between predicted stomatal 1237 conductance rates and what might be observed 1238 through more direct methods, such as sap flux. It 1239 does suggest the hypothesis that the combination of 1240 simulation model and thermal remote sensing data 1241 used in this paper with the particular data set is 1242 sensitive to the same tradeoffs in stomatal regulation 1243 of leaf water potential that is more directly quantifi-1244 able with sap flux. A study that combines ground 1245 validation, including sap flux, combined with thermal 1246 remote sensing data should be able to test this 1247 hypothesis. 1248

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Onion Creek 1985 Julv -dGs/dIn*D* [mmol m⁻² s⁻¹ In(kPa) ⁻¹] August Theoretical 0.6 line 0.8 □ August 0.75 *m /G* s_{ref} [In(kPa)⁻¹] July 0.7 0.65 0.6 0.55 0.5 └─ 0.05 0.055 0.06 R' (m) $G_{\text{Sref}} \pmod{m^2 s^{-1}}$ Fig. 8. These plots show the relationships between stomatal sensitivity to vapor pressure deficit and reference conductance at a vapor pressure

¹²⁷⁰ ¹²⁷² ¹²⁷² ¹²⁷² ¹²⁷² ¹²⁷² ¹²⁷² ¹²⁷² ¹²⁷² ¹²⁷³ ¹²⁷⁴ ¹²⁷⁴ ¹²⁷⁴ ¹²⁷⁴ ¹²⁷⁴ ¹²⁷⁴ ¹²⁷⁵ ¹²⁷⁶ ¹²⁷⁶ ¹²⁷⁶ ¹²⁷⁶ ¹²⁷⁶ ¹²⁷⁷ ¹²⁷⁷ ¹²⁷⁷ ¹²⁷⁸ ¹²⁷⁸ ¹²⁷⁸ ¹²⁷⁸ ¹²⁷⁸ ¹²⁷⁸ ¹²⁷⁹

The results suggest that the four parameters, g_{Smax} , δ , Q_{\min} , and R'_h are sufficient for parameterizing the Jarvis model within RHESSys applied to conifer-covered watersheds. Among hillslope variations in Q_{\min} rooting depths or some similar proxy of available soil moisture appear to explain much of the variation among hillslopes. Once these are accounted for, g_{Smax} and δ parameters can be identified and are nearly equal for all hillslopes. While this is not necessarily representative of individual locations within the watershed, the average values of maximum stomatal conductance obtained are typical for conifer canopies (Running and Coughlan, 1988; Kelliher et al., 1995). This suggests that, at a scale of hillslopes or larger, average canopy parameters are acceptable, at least for the conifer biome. While this approach used to calibrate RHESSys does not resolve within-hillslope details in canopy parameters, it does allow for an assessment of physically meaningful parameter trade-offs. The tradeoffs between canopy parameters indicate a physiological basis for interpreting simulated transpiration. Based on the universal line transpiration rates are reduced either by lowering G_{Sref} or by increasing *m*. To prevent runaway cavitation a plant either needs to have a low set point for its leaf water potential, which requires a high structural integrity of its cell walls (or low vulner-ability to cavitation), or it must safeguard against high leaf water potentials by closing stomata. On one hand, the 'efficiency' associated with a high G_{Sref} comes at a cost, as high $K_{\rm L}$ per unit L is needed to meet the high demand for water, and this makes the plant vulnerable to hydraulic failure (Ewers et al., 2000). On the other hand, the 'safety' associated with a low G_{Sref} means the plant can have a lower K_L per unit L. While this reduces its vulnerability to hydraulic failure when water is limiting or atmospheric demand for water is high it also compromises the photosynthetic capacity when water supply exceeds demand. This has direct implications for carbon gain and plant growth. The simulation model mimics this by either reducing g_{Smax} or increasing δ . A higher g_{Smax} allows a simulated

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plant to take advantage of optimal environmental 1345 conditions to maximize CO2 gain. These differences 1346 in stomatal response to environmental factors have 1347 important implications for land surface process 1348 modeling of water and carbon exchange, from short 1349 to long timescales. The safety versus efficiency 1350 tradeoffs embodied provides a direct physical con-1351 nection between model parameterization and the 1352 physiological functioning of vegetation at watershed 1353 scales. As such, this model could be considered 1354 1355 complimentary to the traditional land surface parameterization schemes based on biome classification 1356 coupled with remote sensing (Running et al., 1995; 1357 Sellers et al., 1996; Zeng et al., 2000). Parameteriza-1358 tion of the variability in canopy physiology at large 1359 scales then amounts to mapping the land surface into 1360 positions along the continuum between safe and 1361 efficient strategies. This could greatly simplify the 1362 task of quantifying surface resistances over large 1363 regions. 1364

A major challenge in parameterizing surface 1365 resistance parameters from remote sensing is the 1366 typical low signal-to-noise ratio. This has lead to the 1367 use of thermal inertia (Anderson et al., 1997; Norman 1368 et al., 2000) and empirical relations between veg-1369 etation vigor and surface temperature (Nemani and 1370 Running, 1989; Carlson et al., 1995) to effectively 1371 separate information from noise. The approach in this 1372 paper is similar in that an empirical relationship 1373 between reference conductance and stomatal sensi-1374 tivity to water loss is used to distinguish between 1375 signal and noise. The approach is similar to that of 1376 Franks and Beven (1997) in that spatial variation in 1377 surface temperature is used to constrain the identifi-1378 cation of model parameters. A physical interpretation 1379 1380 in this paper relies on strong gradients in vegetation 1381 types (grass versus gallery forest). The present paper takes this type of analysis a significant step towards 1382 finding a direct physical interpretation of parameters 1383 in terms of the hydraulic functioning of the veg-1384 etation. This can resolve gradients in physiological 1385 functioning even when taxonomic classification is 1386 highly aggregated, as is typically the case for remote 1387 1388 sensing data. The results suggest a potential refinement of current remote sensing algorithms to include 1389 the universal line describing the regulation of leaf 1390 water potential. This would improve future estimates 1391 1392 of land surface parameters, particularly for forests

where strong gradients in surface temperature do not 1393 necessarily accompany strong gradients in water flux. 1394

6. Conclusions

1398 A Jarvis-type model of stomatal conductance can 1399 be reliably parameterized at a daily time scale in a 1400 meso-scale watershed with a limited amount of 1401 thermal remote sensing imagery. However, parameter 1402 values need to be determined carefully, as there are 1403 numerous compensations among parameters. The key 1404 strength of the approach applied in this paper is that 1405 parameter values are not chosen strictly based on 1406 degree of fit between the simulated latent heat fluxes 1407 and the surface temperatures obtained from the 1408 thermal remote sensing data. Instead, a set of 1409 simulation results that cannot be rejected based on 1410 their information content is used as an ensemble. 1411 Furthermore, the ensemble parameter values are 1412 compared against water-relations theory, which 1413 lends a direct physical interpretation to the estimated 1414 parameters. The results are in agreement with those 1415 obtained in other studies of direct analysis of leaf-1416 level porometry measurements, whole-tree sap flux 1417 measurements, and whole canopy modeling validated 1418 with sap flux measurements. While further work is 1419 needed to assess the reliability of the approach across 1420 a range of biomes, it appears to provide a direct link 1421 between estimating stomatal conductance from 1422 remote sensing data and more direct ground obser-1423 vations of plant water-relations. 1424

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