

Capturing soil-water and groundwater interactions with an iterative feedback coupling scheme: New HYDRUS package for MODFLOW

Jicai Zeng, Jinzhong Yang, Yuanyuan Zha, Liangsheng Shi

State Key Laboratory of Water Resources and Hydropower Engineering Science, Wuhan University, Wuhan 430072, China

5 *Corresponding author:* Yuanyuan Zha (zhayuan87@gmail.com)

Abstract. Accurately capturing the complex soil-water and groundwater interactions is vital for describing the coupling between subsurface/surface/atmospheric systems in regional-scale models. The non-linearity of the Richards' equation for water flow, however, introduces numerical complexity to large unsaturated-saturated modeling systems. An alternative is to use quasi-3D methods with a feedback coupling scheme to practically join sub-models with different properties, such as

10 governing equations, numerical scales, and dimensionalities. In this work, to reduce the non-linearity in the coupling system, two different forms of the Richards' equation are switched according to the soil-water content at each numerical node. A rigorous multi-scale water balance analysis is carried out at the phreatic interface to link the soil water and groundwater models at separated spatial and temporal scales. For problems with dynamic groundwater flow, the non-trivial coupling errors introduced by the saturated lateral fluxes are minimized with a moving-boundary approach. It is shown that the developed

15 iterative feedback coupling scheme results in significant error reduction, and is numerically efficient for capturing drastic flow interactions at the water table, especially with dynamic local groundwater flow. The coupling scheme is developed into a new HYDRUS package for MODFLOW, which is applicable for regional-scale problems.

Key words: Soil-water-groundwater interaction; Multi-scale water balance; Iterative feedback coupling; Regional-scale modeling; HYDRUS package for MODFLOW

Numerical modeling of the soil-water and groundwater interactions has to deal with flow components and governing equations at different scales. This adds significant complexity to model development and calibration. Unsaturated soil water and saturated groundwater flows with similar properties are usually integrated into a whole modeling system. Although physically consistent and numerically rigorous, methods involving the 3D Richards' equation (*RE*, (Richards, 1931)) tend to be computationally expensive and numerically unstable due to the large non-linearity and the demand for dense discretization (Kumar et al., 2009; Maxwell and Miller, 2005; Panday and Huyakorn, 2004; Thoms et al., 2006; Zha et al., 2013a), especially for problems with multi-scale properties. In this work, parsimonious approaches, which appear in different governing equations and coupling schemes, are developed for modeling the soil-water and groundwater interactions at regional scales.

Simplifying the soil-water flow details into upper flux boundaries has been widely used to simulate large-scale saturated flow dynamics, such as MODFLOW package and its variants (Langevin et al., 2017; Leake and Claar, 1999; McDonald and Harbaugh, 1988; Niswonger et al., 2011; Panday et al., 2013; Zeng et al., 2017). At local scale in contrast, the unsaturated flow processes are usually approximated with reasonable simplifications and assumptions in the Richards' equation (Bailey et al., 2013; Liu et al., 2016; Paulus et al., 2013; Šimůnek et al., 2009; van Dam et al., 2008; Yakirevich et al., 1998; Zha et al., 2013b).

The original Richards' equation, also the *mixed-form RE*, takes pressure head (h) as the driving force variable, while soil moisture content (θ) as the mass accumulation variable (Krabbenhøft, 2007). To solve the *mixed-form RE*, either h or θ , or a *switching* of both, is assigned as the primary variable. The *h-form RE* is widely employed for unsaturated-saturated flow simulation, especially in heterogeneous soils, such as the HYDRUS package (Šimůnek et al., 2016). Significant improvement in mass conservation has been achieved with Celia's modification (Celia et al., 1990). Then, efforts were made to combine the advantages in the *original* and the *Celia-format h-form REs* by switching their storage terms (Hao et al., 2005; Zadeh, 2011). However, these models still suffer from high computational cost and low numerical robustness when dealing with rapidly changing atmospheric boundary conditions (Crevoisier et al., 2009; Zha et al., 2017). The *θ -form RE*, addressing the above problems, is inherently mass conservative and less non-linear in the averaged nodal hydraulic diffusivity when the soil is dry (Warrick, 1991; Zha et al., 2013b). However, the *θ -form RE* is not applicable for saturated and heterogeneous soils (Crevoisier et al., 2009; Zha et al., 2013b). In this work, to take advantages of both forms of *RE*, the governing equations, rather than primary variables (Diersch and Perrochet, 1999; Forsyth et al., 1995; Zha et al., 2013a), are switched at each node according to its saturation degree.

For regional problems, the vadose zone is usually conceptualized into paralleled soil columns without lateral connections. The resulting quasi-3D coupling scheme (Kuznetsov et al., 2012; Seo et al., 2007; Xu et al., 2012; Zhu et al., 2012) significantly

50 reduces the dimensionality and complexity. According to how the messages are transferred across the coupling interface, the quasi-3D methods are categorized into (1) the fully coupling scheme, which simultaneously builds the nodal hydraulic connections of sub-models at both sides and implicitly solves the assembled matrices; (2) the one-way coupling scheme, which delivers the soil-water model solutions onto the groundwater model without feedback mechanism; and (3) the feedback (or two-way) coupling scheme, which explicitly exchanges the head/flux solutions in vicinity of the interface nodes.

55 The fully coupling scheme (Gunduz and Aral, 2005; Zhu et al., 2012) is numerically rigorous but tends to increase the computational burden under practical conditions. For example, the potentially conditional diagonal dominance causes non-convergence for the iterative solvers (Edwards, 1996). Owing to high non-linearity in the soil-water sub-models, the assembled matrices can only be solved with unified small time-steps, which adds to the computational expense. The one-way coupling scheme, as adopted by the UZF1 package for MODFLOW (Grygoruk et al., 2014; Niswonger et al., 2006), as well as the free
60 drainage mode in SWAP package for MODFLOW (Xu et al., 2012), assumes that the water table depth is of minor influence on flow interactions at the phreatic interface, and is thus problem specific.

The feedback coupling methods, in contrast, are widely used (Kuznetsov et al., 2012; Seo et al., 2007; Shen and Phanikumar, 2010; Stoppelenbrug et al., 2005; Xie et al., 2012; Xu et al., 2012) as a compromise of numerical accuracy and computational cost. In a feedback coupling scheme, the soil-water and groundwater sub-models can be built with governing equations and
65 numerical schemes at different scales. For flow processes with multi-scale components, such as boundary geometries, parameter heterogeneities, and hydrologic stresses, the scale-separation strategy can be implemented easily. Although the feedback coupling method is numerically more rigorous than a one-way coupling method, and tends to reduce the inconsistency of head/flux interfacial boundaries, some concerns arise.

The first concern is the numerical efficiency of the iterative/non-iterative feedback coupling methods. The non-iterative
70 approach (Twarakavi et al., 2008; Xu et al., 2012) usually leads to significant error accumulation when dealing with dynamically fluctuating water table, especially with large time-step sizes. The iterative methods in contrast (Kuznetsov et al., 2012; Stoppelenbrug et al., 2005; Xie et al., 2012), by converging the head/flux solutions at the coupling interface, are numerically rigorous but computationally expensive, especially when solving the coupled sub-models with a unified time-stepping scheme (Kuznetsov et al., 2012). Good balance between cost and effect is needed to maintain practical utility of the
75 iterative feedback coupling scheme.

The second concern lies in the scale-mismatching problem. For groundwater models (Harbaugh et al., 2017; Langevin et al., 2017; Lin et al., 2010; McDonald and Harbaugh, 1988), the specific yield at the phreatic surface is usually represented by a simple large-scale parameter; while for soil-water models (Niswonger et al., 2006; Šimůnek et al., 2009; Thoms et al., 2006), the small-scale phreatic water release is influenced by the water table depth and the unsaturated soil moisture profile (Dettmann

80 and Bechtold, 2016; Nachabe, 2002). Delivering small-scale solutions of the soil-water models onto the large-scale interfacial boundary of the groundwater model, as well as maintaining the global mass balance, usually introduce significant non-linearity to the entire coupling system (Stoppelenburg et al., 2005). Conditioned by this, the mismatch of numerical scales in the coupled sub-models causes significant coupling errors and instability. A multi-scale water balance analysis at the phreatic surface helps to relieve such difficulties.

85 The third concern is the non-trivial lateral fluxes between the saturated regions controlled by the vertical soil columns, which are usually not considered in previous study (Seo et al., 2007; Xu et al., 2012). Though rigorous water balance analysis is conducted to address such inadequacy (Shen and Phanikumar, 2010), the lateral fluxes solved with a 2D groundwater model usually require additional efforts to build water budget equations in each sub-division represented by the soil columns. A moving boundary strategy helps to avoid the saturated lateral flow in the groundwater body.

90 In this work, the h - and θ -form of the 1D RE are switched at equation level to obtain a new HYDRUS package. To handle three of the aforementioned concerns, a multi-scale water balance analysis is carried out at the phreatic surface to conserve head/flux consistent at the coupling interface. An iterative feedback coupling scheme is developed for linking the unsaturated and saturated flow models at disparate scales. The saturated lateral fluxes between the soil columns are fully removed from the interfacial water balance equation, making it a moving-interface coupling framework. The head solution of MODFLOW-2005
95 (Harbaugh et al., 2017; Langevin et al., 2017) and flux solution of HYDRUS1D (Šimůnek et al., 2009), are relaxed to meet consistency at the phreatic surface.

In this paper, the governing equations at different scales, the multi-scale water balance analysis at the phreatic surface, and the iterative feedback coupling scheme for solving the whole system, are presented in Section 2. Synthetic numerical experiments are described in Section 3. Numerical performance of the developed model is investigated in Section 4. Conclusions are drawn
100 in Section 5.

2 Methodology

To address the aforementioned first concern, governing equations for subsurface flow are given at different levels of complexity (section 2.1); numerical solution of these equations are presented (section 2.2); nonlinearity in the soil-water sub-models are reduced by a generalized switching scheme that chooses appropriate forms of the Richards' equation (RE) according to the
105 hydraulic conditions at each numerical node (section 2.3); then, an iterative feedback coupling scheme is developed to solve the soil-water and groundwater models at independent scales (section 2.4). As for the second concern, a multi-scale water balance analysis is conducted to deal with the scale-mismatching problem at the phreatic surface (section 2.5). To cope with the third concern, a moving Dirichlet boundary at the groundwater table is assigned to the soil water sub-models (see Appendix

A.1); the Neumann upper boundary for the saturated model is provided in Appendix A.2; the relaxed iterative feedback process
 110 is presented in Appendix A.3.

2.1 Governing equations

The mass conservation equation for unsaturated-saturated flow is given by:

$$\frac{\partial \theta}{\partial t} + \beta \mu_s \frac{\partial h}{\partial t} = (C + \beta \mu_s) \frac{\partial h}{\partial t} = -\nabla \cdot \mathbf{q} \quad (1)$$

where t is time [T]; θ [L^3L^{-3}] is volumetric moisture content; h [L] is pressure head; β is one for saturated region while zero
 115 for the unsaturated region; C [L^{-1}] is the soil water capacity ($C = \partial\theta/\partial h$) for unsaturated region, while zero for saturated region;
 μ_s [L^{-1}] is specific elastic storage; \mathbf{q} [LT^{-1}] is Darcian flux calculated by:

$$\mathbf{q} = -K\nabla H \quad (2)$$

where K [LT^{-1}] is the hydraulic conductivity, $K = K(\theta)$; H [L] is the potentiometric head, $H = h + z$, in which z is the vertical
 120 location with coordinate positive upward. Combining Eqns. (1) and (2) results in the governing equation for groundwater
 flow

$$\mu_s \frac{\partial H}{\partial t} = \frac{\partial}{\partial x} \left(K \frac{\partial H}{\partial x} \right) + \frac{\partial}{\partial y} \left(K \frac{\partial H}{\partial y} \right) + \frac{\partial}{\partial z} \left(K \frac{\partial H}{\partial z} \right) \quad (3)$$

With the assumption that the horizontal unsaturated flows are negligible, the regional vadose zone is usually represented by
 an assembly of paralleled soil columns. The generalized 1D *RE* is represented by a switchable format,

$$\hat{C} \frac{\partial \psi}{\partial t} = \frac{\partial}{\partial z} \left(\hat{K} \left(\frac{\partial \psi}{\partial z} + 1 \right) \right) \quad (4)$$

125 where ψ is the primary variable. For an *h-form RE*, $\psi = h$, $\hat{C} = C$, and $\hat{K} = K$; while for a *θ -form RE*, $\psi = \theta$, $\hat{C} = 1$,
 $\hat{K} = D$, where D [L^2T^{-1}] is the hydraulic diffusivity, $D = K/C$.

2.2 Numerical approximation

The governing equation for the saturated zone (Eqn. (3)) is spatially and temporally approximated in the same form with the
 MODFLOW-2005 model (Harbaugh et al., 2017; Langevin et al., 2017). Celia's modification (Celia et al., 1990; Šimůnek et
 130 al., 2009) is applied to the *h-form 1D RE* for temporal approximation. Both forms of *RE* are handled with a temporally
 backward finite difference discretization (Zha et al., 2013b, 2017). Each sub-model is solved by a Picard iteration scheme,
 which is widely used in some popular codes/software packages (van Dam et al., 2008; Šimůnek et al., 2016).

The spatial discretization of Eqn. (4), as well as the water balance analysis for each node, are based on the nodal flux in
 element $i+1/2$ (bounded by nodes i and $i+1$), which is

$$q_{i+1/2}^{\psi} = -\frac{\hat{K}_{i+1/2}^{j+1,k}}{\Delta z_{i+1/2}} (\psi_{i+1}^{j+1,k+1} - \psi_i^{j+1,k+1}) - K_{i+1/2}^{j+1,k} + \varepsilon_{i+1/2}^{j+1,k} \quad (5)$$

where the superscripts j and k are the levels of time and inner iteration; the subscript i (or $i+1/2$) is the number of node (or element); $\Delta z_{i+1/2}$ is the length of the element $i+1/2$, $\Delta z_{i+1/2} = (z_{i+1} - z_i)$. When a soil interface exists at node i for example, the soil moisture contents in elements $i-1/2$ and $i+1/2$ are discontinuous at node i , thus dissatisfying the θ -form RE. To address this problem, the correction term $\varepsilon_{i+1/2}^{j+1,k}$, suggested by Zha et al. (2013b), is employed to handle the heterogeneous interface at nodes i and $i+1$,

$$\varepsilon_{i+1/2}^{j+1,k} = \frac{\hat{K}_{i+1/2}^{j+1,k}}{\Delta z_{i+1/2}} \left(\psi_{i+1}^{j+1,k} - \psi_i^{j+1,k} - \bar{\psi}_{i+1}^{j+1,k} + \bar{\psi}_{i,\Omega}^{j+1,k} \right) \quad (6)$$

where $\bar{\psi}_{i+1}^{j+1,k}$ and $\bar{\psi}_i^{j+1,k}$ are the continuously distributed ψ within element $i+1/2$, i.e., between the vertices i and $i+1$.

When $\psi = h$, or when $\psi = \theta$ but no heterogeneity occurs, we get $\psi_{i+1}^{j+1,k} = \bar{\psi}_{i+1}^{j+1,k}$ and $\psi_i^{j+1,k} = \bar{\psi}_i^{j+1,k}$, so $\varepsilon_{i+1/2}^{j+1,k} = 0$. When $\psi = \theta$, with soil interfaces at node i or $i+1$, $\bar{\psi}_{i+1}^{j+1,k} = \theta(h_{i+1}^{j+1,k}, \mathbf{p}_{i+1/2})$ and $\bar{\psi}_i^{j+1,k} = \theta(h_i^{j+1,k}, \mathbf{p}_{i+1/2})$. It is obvious that

$\psi_i^{j+1,k} \neq \bar{\psi}_i^{j+1,k}$ (or $\psi_{i+1}^{j+1,k} \neq \bar{\psi}_{i+1}^{j+1,k}$), so $\varepsilon_{i+1/2}^{j+1,k} \neq 0$.

Hereinafter, $\mathbf{P}_{i+1/2}$ represents the soil parameters in element $i+1/2$. For example, in van Genuchten model (van Genuchten, 1980), $\mathbf{P}_{i+1/2} = (\theta_r, \theta_s, n, m, \alpha, k_s)$, where θ_r [L^3L^{-3}] and θ_s [L^3L^{-3}] are the residual and saturated soil moisture contents; α [L^{-1}], n , and m are the pore-size distribution parameters, $m = 1-1/n$; k_s [LT^{-1}] is the saturated hydraulic conductivity.

2.3 Switching the Richards' equation

Due to lower non-linearity of hydraulic diffusivity (D) for dry soils (Zha et al., 2013b) and the avoidance of mass balance error by removing the soil water capacity in the storage term, the θ -form RE is more robust than the h -form RE, especially when dealing with rapidly changing atmospheric boundary conditions (Zeng et al., 2018). In our work, the h - and θ -form REs are switched at each node according to its effective saturation, Se . The resulting hybrid matrix equation set is solved by Picard iteration. The empirical effective saturation for doing switching varies with soil type and is suggested to be $Se^{crit} = 0.4-0.9$, the state when both the h - and θ -form REs are stable and efficient. When $Se \geq Se^{crit}$, the soil moisture is closer to saturation, so the h -form RE is chosen as the governing equation; otherwise, when it undergoes dry soil condition, the θ -form RE is preferred.

For element $i+1/2$, when the governing equations for nodes i and $i+1$ are identical, the spatial approximation of nodal flux is given by Eqn. (5). When the governing equations differ at nodes i and $i+1$, a switched element is produced. Take $\psi_i = \theta_i$ and $\psi_{i+1} = h_{i+1}$ for example, the nodal fluxes calculated by Eqn. (5) for different forms of RE have to be carefully handled

by substituting $\theta_{i+1}^{j+1,k+1}$ with $\theta_{i+1}^{j+1,k}$, while $h_i^{j+1,k+1}$ is replaced by $h_i^{j+1,k}$. When $\psi_i = h_i$ and $\psi_{i+1} = \theta_{i+1}$, in contrast, $h_{i+1}^{j+1,k+1}$ is replaced by $h_{i+1}^{j+1,k}$, while $\theta_i^{j+1,k+1}$ is replaced by $\theta_i^{j+1,k}$. The resulting equivalent nodal fluxes $q_{i+1/2}^h$ and $q_{i+1/2}^\theta$ are then weighted to obtain an approximation by

$$q_{i+1/2} = (1-\omega)q_{i+1/2}^\theta + \omega \cdot q_{i+1/2}^h \quad (7)$$

where ω is the weighting factor, $0 \leq \omega \leq 1$. In our work, $\omega = 0.5$ is applied to implicitly maintain the unknown variables of

165 both $h_{i+1}^{j+1,k+1}$ and $\theta_i^{j+1,k+1}$. Specifically, when $\omega = 1$, the *h-form RE* is used at nodes i and $i+1$; when $\omega = 0$, the *θ -form RE* is employed instead. A detailed study on doing switching of *RE* between two ends of the soil moisture condition, as well as the description of the numerical formation can be found in Zeng et al. (2018).

Note that the equation switching method takes full advantages of the *θ -* and *h-form REs*, which is different from the traditional primary variable switching schemes (Diersch and Perrochet, 1999; Forsyth et al., 1995; Zha et al., 2013a). In our work, the
170 *switching-RE* approach is incorporated into a new HYDRUS package.

2.4 Iterative feedback coupling scheme

The Dirichlet and Neumann boundaries are iteratively transferred across the phreatic interface. The groundwater head solution serves as the head-specified lower boundary of the soil columns; while the unsaturated solution is converted into the flux-specified upper boundary of the groundwater model. Due to moderate variation of the groundwater flow, the predicted water-
175 table solution is usually adopted in advance as Dirichlet lower boundary of the fine-scale soil-water flow models (Seo et al., 2007; Shen and Phanikumar, 2010; Xu et al., 2012), which then in sequence provides the Neumann upper boundary for successively solving the coarse-scale groundwater flow model. **Appendix A.1** provides the method for a moving Dirichlet lower boundary, while **Appendix A.2** presents the Neumann upper boundary for the 3D groundwater model. In **Appendix A.3**, the relaxed iterative feedback coupling scheme is used to solve the unsaturated/saturated sub-models at two sides of the
180 coupling interface.

2.5 Multi-scale water balance analysis

Coupling models at different scales requires consistency in their spatial and temporal scales at the interface (Downer and Ogden, 2004; Rybak et al., 2015). Space- and time-splitting strategy (see **Figure 1**) are adopted to separate sub-models at different scales. That is, the soil water models are established by $\Delta z = 10^{-3}$ m- 10^0 m, and $\Delta t = 10^{-5}$ d- 10^0 d; while for the
185 saturated model, the grid sizes are $\Delta x = 10^0$ m- 10^3 m, and time-step sizes are $\Delta t = 10^0$ d- 10^1 d. Water balance at one side of the interface is conserved by scale matching of boundary conditions provided by the sub-model on the other side. For unsaturated flow, the Richards' equation requires fine discretization of space and time (Miller et al., 2006; Vogel and Ippisch, 2008); while for saturated flow, coarse spatial and temporal grids produce adequate solutions at large scale (Mehl and Hill, 2004; Zeng et al., 2017). To approximate the upper boundary flux of the groundwater flow model, a multi-scale water balance analysis is
190 conducted within each step of the large-scale saturated flow model. At small spatial and temporal scales, e.g., within a macro time step $\Delta T = T^{j+1} - T^j$ and at a local area of interest (with thickness of $\bar{M} = z_s - z_0$, where z_s and z_0 are defined in **Appendix A.2**), the specific storage term in Eqn. (1) is vertically integrated into a transient one-dimensional expression (Dettmann and Bechtold, 2016),

$$\tilde{S}_y = \left[w(T^{J+1}) - w(T^J) + \theta_s \cdot \Delta z_t \right] / \Delta z_t + \mu_s \cdot \bar{M} \quad (8)$$

195 where w [L] is the amount of unsaturated water in the moving balancing domain, see **Figure 2b**, $w(t) = \int_{z_t(t)}^{z_s} \theta(t, z) dz$; $\Delta z_t = \sum_{j=1}^N dz_t^j = z_t(T^{J+1}) - z_t(T^J)$ is the total fluctuation of the phreatic surface during $\Delta T = \sum_{j=1}^N dt^j = T^{J+1} - T^J$; θ_s is the saturated soil water content. Approaching a transient state at time t , the water balance in a moving water balancing domain (see $z \in [z_t, z_s]$ in **Figure 2b**) during a small-scale time step dt (defined in **Figure 1b**) is given by

$$[q_{top} + l \cdot dz_t / 2 - q_{bot}] \cdot dt = w(t) - w(t - dt) + \theta_s \cdot dz_t \quad (9)$$

200 where $q_{top}(t)$ and $q_{bot}(t)$ [LT⁻¹] are the nodal fluxes into and out of the moving balancing domain at a fixed top boundary (z_s) and a moving bottom boundary ($z_b = \min(z_t(t), z_t(t - dt))$), $q_{top} = K(h) \cdot \partial(h + z) / \partial z|_{z=z_s}$, $q_{bot} = K(h) \cdot \partial(h + z) / \partial z|_{z=z_b}$ (positive into the balancing domain and negative outside); $dz_t = z_t(t) - z_t(t - dt)$ is the transient fluctuation of the phreatic surface during dt ; l [T⁻¹] is the saturated lateral flux into the balancing domain at time t , see **Figure 2b**. Taking Γ as the lateral boundary of a

sub-domain, the lateral flux $l = \iiint_{x,y,z \in \Omega} \left[\frac{\partial}{\partial x} \left(K \frac{\partial H}{\partial x} \right) + \frac{\partial}{\partial y} \left(K \frac{\partial H}{\partial y} \right) \right] dx dy dz / \iiint_{x,y,z \in \Omega} dx dy dz$ is supposed to be constant during ΔT ;

205 Ω is the volume of the saturated domain controlled by a soil column, which is horizontally projected into Π . Temporally integrating Eqn. (9) from time T^J to T^{J+1} produces

$$R_{top} + \varepsilon_l - R_{bot} = w(T^{J+1}) - w(T^J) + \theta_s \cdot \Delta z_t \quad (10)$$

where R_{top} [L] is the cumulative water flux at z_s , $R_{top} = \int_{T^J}^{T^{J+1}} q_{top}(t) dt$, note that R_{top} equals F_{top} in Eqn. (A3) (**Appendix A.2**);

R_{bot} [L] is the cumulative water flux out of the moving balancing domain, $R_{bot} = \int_{T^J}^{T^{J+1}} q_{bot}(t) dt$; ε_l [L] is the cumulative

210 lateral input water into the moving balancing domain,

$$\varepsilon_l = \frac{1}{2} l \cdot \sum_{j=1}^N dt^j dz_t^j \quad \varepsilon_l' = \frac{1}{2} l \cdot \Delta T \cdot \Delta z_t \quad (11)$$

where N is the number of time steps for the small-scale soil-water model within a macro time step ΔT ; and ε_l' is the non-trivial saturated later flux produced by a stationary boundary method (Seo et al., 2007; Xu et al., 2012). By taking R_{top} as the specific recharge at z_s , the small-scale specific yield \tilde{S}_y is derived from Eqns. (8) and (10) as

$$215 \quad \tilde{S}_y = (R_{top} + \varepsilon_l - R_{bot}) / \Delta z_t + \mu_s \cdot \bar{M} \quad (12)$$

Suppose z_t is linearly fluctuating in time, i.e., $z_t = a \cdot t + b$, (where a and b are constants), we get the water table change during a small-scale step (dt) by $dz_t = a \cdot dt$, thus, $\varepsilon_l = \alpha(dt^2)$, which means linearly refining the local time-step size (dt) in the soil water model brings about at least quadratic approximation of ε_l towards zero. Thus ε_l can be neglected from the small-scale mass balance analysis. In the developed model, the large-scale specific yield, \bar{S}_y in Eqn. (A3), represents the water release in the

220 phreatic aquifer; while the small-scale \tilde{S}_y in Eqn. (12), denotes the dynamically changing water yield caused by the fluctuation of the water table. The upper boundary flux F_{top} in the phreatic flow equation (Eqn. (A3)) is therefore corrected to

$$F_{top} = [R_{top} + (\bar{S}_y - \tilde{S}_y)\Delta z_t] / \Delta T \quad (13)$$

225 Differing from previous studies (Seo et al., 2007; Shen and Phanikumar, 2010; Xu et al., 2012), a scale-separation strategy is employed in Eqn. (13). The specific yields at two different scales are explicitly linked in F_{top} . The large-scale properties in the groundwater model (MODFLOW) are thus fully maintained.

3 Numerical experiments

In this section, a range of 1D, 2D, 3D, and regional numerical test cases are presented. The 1D tests are benchmarked by the globally refined solutions from the HYDRUS1D code (Šimůnek et al., 2009). The 2D/3D “truth” solutions are obtained from the fully-3D unsaturated-saturated flow model VSF (Thoms et al., 2006). At regional scale, a synthetic case study suggested by Twarakavi et al. (2008) is reproduced. The codes are run on a 16 GB RAM, 3.6 GHz Intel Core (i3-4160) based personal computer. A maximal number of feedback iteration is set at 20. Soil parameters for the van Genuchten model (van Genuchten, 1980) are given in Table 1. The root mean square error (RMSE) of the solution ψ at time t is given by

$$\text{RMSE}(\psi, t) = \left\{ \frac{1}{N} \sum_{i=1}^N (\psi_i^{ref}(\mathbf{x}, t) - \psi_i(\mathbf{x}, t))^2 \right\}^{1/2} \quad (14)$$

230 where ψ is the numerical solution of either pressure head or water content; ψ^{ref} is the corresponding reference solution; Subscript i is the number of nodes, $i = 1, 2, \dots, N$.

3.1 Case 1: Rapidly changing atmospheric boundaries

The 1D case is used to investigate the benefit brought by switching the Richards’ equation in the unsaturated zone. A soil column is initialized with hydrostatic water-table depth of 800 cm. That is, $h(t = 0, z) = 200 - z$ cm, with $z = 0$ at the bottom, and $z = 1,000$ cm on the top. The lower boundary is set non-flux to avoid the extra computational burden caused by variation of the groundwater model. Two scenarios from literature are reproduced with rapidly changing upper boundaries, as well as extreme flow interactions between the unsaturated and saturated zones.

240 Miller et al.’s problem (Miller et al., 1998) is reproduced in scenario 1. A dry-sandy soil column (see soil #1 in Table 1) experiences a large constant flux infiltration at the soil surface of $q_{top} = 30$ cm/d which ceases at $t = 4$ d.

In scenario 2, Hills et al.’s problem (Hills et al., 1989) is considered. The soils #2 and #3 from Table 1 are alternatively layered with a thickness of 20 cm within the first 80-cm depth. Below 80 cm ($z = 0-920$ cm) is soil #2 with non-flux bottom boundary.

The atmospheric upper boundary conditions, rainfall and evaporation change rapidly with time (see **Figure 3**), over 365 days. The coupled unsaturated model is discretized into a fine grid with $\Delta z = 1$ cm, while the saturated model is discretized into two

layers with thickness of 500 cm. The impact of different numbers of feedback iteration, closure criteria, as well as different forms of 1D Richards' equation, are investigated. Solutions obtained from the HYDRUS1D model with $\Delta z = 1$ cm, and $\Delta t =$
250 0.05 d are taken as the "truth".

3.2 Case 2: Dynamic Groundwater flow

A 2D case is analyzed with sharp groundwater flow (see **Figure 4**). To minimize the unsaturated lateral flow, the soil surface is set with non-flux boundary. The bottom and lateral boundaries are also non-flux. Two pumping stresses are applied to the cross-sectional field with $x \times z = 5,000$ cm \times 1,000 cm. Well #1 is located at $x = 2,500$ cm, with pumping screen at $z = 0-200$
255 cm; while well #2 is at $x = 5,000$ cm, with pumping screen of $z = 0-200$ cm. Pumping rates for wells #1 and #2 respectively are 2×10^4 cm²/d and 1×10^4 cm²/d per width unit. The initial hydrostatic head of the cross-section is $h_0(x, z) = 700$ cm. Soil #4 in Table 1 fills the entire cross-section. The total simulation lasts 50 days. For the coupled saturated sub-model, as well as the reference model (VSF (Thoms et al., 2006)), the cross-section is discretized horizontally into uniform segments with width $\Delta x = 50$ cm, while vertically (bottom-up) refined into segments with thickness $\Delta z = 200$ cm_($\times 1$), 100 cm_($\times 2$), 50 cm_($\times 2$), 25 cm_($\times 2$),
260 12.5 cm_($\times 4$), and 5 cm_($\times 200$), where the subscripts hereinafter ($\times N$) are the numbers of discretized segments. The 1D soil water models are discretized with segmental thickness of $\Delta z = 1$ cm. The fully-2D unsaturated-saturated solutions from VSF model are taken as the "truth".

3.3 Case 3: Pumping and irrigation

Case 3 is used to investigate the efficiency and applicability of a quasi-3D coupling model in comparison of the fully-3D
265 approaches. A phreatic aquifer with $x \times y \times z = 1,000$ m \times 1,000 m \times 20 m is stressed by constant irrigation and pumping wells. The infiltration rate is 3 mm/d in $(x, y) = (0-440$ m, 560 m-1,000 m), while 5 mm/d in $(x, y) = (560$ m-1,000 m, 0-440 m). Screens for three of the pumping wells locate at $(x, y, z) = (220$ m, 220 m, 5-10 m), (500 m, 500 m, 5-10 m), and (780 m, 780 m, 5-10 m). The pumping rates are constant at 30 m³/d. The initial hydrostatic head of the aquifer is 18 m. Around and below the aquifer are non-flux boundaries. The aquifer is horizontally discretized with $\Delta x = \Delta y = 40$ m for the coupled saturated
270 model, as well as for the VSF model for obtaining the "truth" solution. The top-down thicknesses of the fully-3D grid are $\Delta z = 0.10$ m_($\times 30$), 0.4 m_($\times 5$), 1 m_($\times 5$), and 2 m_($\times 5$). For the 1D soil columns, $\Delta z = 0.1$ m_($\times 30$), and 0.4 m_($\times 5$), which means no soil column reaches the bottom. Different numbers of the sub-zones represented by soil columns, as well as their different geometries, are given in **Figure 5**. The soil parameters for a sandy loam (soil #5) are given in Table 1. Total simulation lasts 60 days.

3.4 Case 4: Synthetic regional case study

275 A hypothetical test case from literature (Niswonger et al., 2006; Prudic et al., 2004; Twarakavi et al., 2008) for large-scale

simulation is reproduced here. The overall alluvial basin is divided into uniform grids with $\Delta x = \Delta y = 1,524$ m. The coupled saturated model is conceptualized into a single layer. The initial head, as well as the elevations of land surface and bedrock, are presented in **Figure 6a**, **b**, and **c**. The precipitation, evaporation, and pumping rates for 12 stress periods, each lasted 1/12 of 365 days, are given in Table 2. The infiltration factors (see **Figure 6d**) are used to approximate the spatial variability of precipitation. The initial head in the vadose zone is set with hydrostatic status. Twenty soil columns, coinciding with the sub-zones in **Figure 6d**, are discretized separately with a range of gradually refined segments with thickness (Δz) from 30.48 cm, to 0.3048 cm (bottom-up). Comparative analysis is conducted with the solutions obtained from the original HYDRUS package for MODFLOW (taken as HPM for short) (Seo et al., 2007).

4 Results and discussion

4.1 Reducing the complexity of a feedback coupling system

The numerical difficulty in a coupled unsaturated-saturated flow system originates from the non-linearity of the soil-water models, heterogeneity of the parameters, as well as the variability of the hydrologic stresses (Krabbenhøft, 2007; Zha et al., 2017). In our work, the overall complexity of an iteratively coupled quasi-3D model could be lowered by (1) taking full advantages of the *h*- and *θ*-form REs to reduce the nonlinearity in the soil-water models, and (2) smoothing the variability in the exchanged interfacial messages.

Two scenarios in case 1 were selected to address the first point. Sudden infiltration into a dry-sandy soil, and the rapidly altering atmospheric upper boundaries, were tested to illustrate the importance of applying a *switching-form RE* for lowering the non-linearity in the soil-water models. To evaluate the benefits brought by a *switching-form RE*, the numerical stability was first considered, as shown in **Figure 7**. The coupled model in our work was tested with *h-form* and *switching-form REs*.

Compared with the HYDRUS1D model (also based on an *h-form RE*), the *switching-form* method was numerically more robust, i.e., with larger minimal time-step sizes (Δt_{min}) and less computational cost, where minimal time-step size was acceptable 10^{-3} d for convergence. Notably at the beginning of the sudden infiltration into a dry-sandy soil, in **Figure 7a**, the Δt_{min} for a switching method was 10^{-3} d, even at early infiltration times, while for the *h-form* methods, including HYDRUS1D and the coupled *h-form* method, Δt_{min} was constrained to 10^{-8} d before reaching a painstaking convergence. In

Figure 8, the soil water content solution by the coupled *switching-form* method and the HYDRUS1D method (taken as the “truth”) were compared at depth of 0, 50 cm, and 200 cm. The RMSEs of the soil moisture solutions (θ) at three different depths are respectively 0.0189, 0.0032, and 0.0013. To finish the calculation, the coupled *switching-form RE* method took 17 seconds, while it was 41 seconds for the HYDRUS code. When solving the same problem, the matrix equation set was solved 4,903 times with the switching scheme, while 10,925 times for the HYDRUS1D code. The switched governing equations

305 contributes to cutting the computational cost by half for problems with rapidly changing upper boundary conditions. Here, the threshold for choosing an appropriate form of RE was non-sensitive to the numerical efficiency. A wide range of $Se^{crit} \in [0.3, 0.9]$ was suggested according substantial trial-and-error tests.

Reducing the complexity of a coupling system can also be attained by smoothing the exchanged information in space and time. As suggested by Stoppelenburg et al. (2005), a time-varying specific yield calculated by the small-scale soil-water models, 310 \tilde{S}_y in Eqn. (12), introduced significant variability to the large-scale groundwater model, thus caused extra iterations. A large-scale \bar{S}_y reduced the non-linearity of the storage term in the groundwater equation. In case 1, using an \bar{S}_y of 0.1-0.2 in the groundwater model produced best numerical stability for the sandy soil with dramatically uprising water table. With a large-scale \bar{S}_y , the non-linearity introduced by the small-scale soil-water models could be quickly smoothed, as shown in Eqn. (12).

315 4.2 Multi-scale water balance analysis

The traditional non-iterative feedback coupling methods cannot maintain sound mass balance near the phreatic surface, especially for problems with drastic flow interactions.

One reason is that, to launch a new step of a sub-model at either side of the phreatic interface, the non-iterative feedback methods usually employed a predicted interfacial boundary without correction, which inevitably introduced coupling errors.

320 In traditional non-iterative methods (Seo et al., 2007; Xu et al., 2012), such shortcomings could be alleviated by refining the macro time step size (ΔT). However, the Dirichlet head predicted for the soil columns with a stepwise extension method (see **Figure 2a**), was easy to implement but tended to suffer from significant coupling error. In this work, we proposed a linear extrapolation method for the lower boundary head prediction for the soil water models, see Eqn. (A2). Here, we used Niter to indicate the maximal number of feedback iteration. Compared with a traditional stepwise method, the solution obtained by a 325 linear method, either iteratively (with Niter = 3) or non-iteratively (Niter = 0), was easier to approach the truth, see **Figure 9**. Even with refined macro time step sizes (ΔT from 0.2 d to 0.005 d), the stepwise method made a thorough effort to minimize the coupling errors. Notably, three feedback iterations (Niter = 3) were sufficient to reduce the coupling error significantly. Such a one-dimensional case with constant upper boundary flux, avoiding interference from lateral fluxes, illustrated the importance of a temporal scale-matching analysis for coupling the soil-water and groundwater models.

330 The other factor contributing to the coupling errors in the traditional method lies in neglecting the saturated lateral flux between adjacent soil columns (Seo et al., 2007; Stoppelenbrug et al., 2005; Xu et al., 2012). In practical applications, the fluxes in and out of the saturated parts of the soil columns differ, which adds to the complexity of the coupling scheme. Although a strict water balance equation is established (Shen and Phanikumar, 2010), the concern centers on the spatial scale-mismatching

problem. That is, when the coarse-grid groundwater flow solutions are converted into the vertically distributed fine-scale source/sink terms for the soil columns, an extra down-scaling approach is needed to ensure their accuracy. Here we carried out a multi-scale water balance analysis above the phreatic surface. The fine-scale saturated lateral flows were thus excluded from Eqn. (10). The benefits of the moving-boundary approach, can be seen in case 2 which produces significant saturated lateral flux. We carried out a comparative analysis against the traditional stationary-boundary methods (Seo et al., 2007; Xu et al., 2012). The 2D solution of VSF was taken as the “truth”. **Figure 10** presents the effectiveness of the moving-boundary method. Five stationary soil columns with three different lengths ($L = 1,000$ cm, 500 cm, and 300 cm) were compared with an adaptively moving soil column within the iterative feedback coupling scheme. The cross-sectional RMSE of the phreatic surface and the head at bottom layer ($z = 0$), are presented in **Figure 10a** and b. The soil columns with bottom nodes fixed deeply into the aquifer, instead of moving with the phreatic surface, introduced large coupling errors. This was caused by the non-trivial saturated lateral fluxes between the adjacent soil columns. With a traditional stationary-boundary method, such problems can be alleviated by avoiding large saturated lateral fluxes between the soil columns. However, for some spatiotemporally varying local events in a regional aquifer (e.g., pumping or flooding irrigation), such problems increased the burden for sub-zone partitioning. A moving-boundary method instead, was numerically more efficient for minimizing the size of the matrix equation and reducing the coupling errors.

4.3 Regulating the feedback iterations

In coupling two complicated modeling system, a common agreement has been reached that, feedback coupling, either iteratively (Markstrom et al., 2008; Mehl and Hill, 2013; Stoppelenbrug et al., 2005; Xie et al., 2012) or non-iteratively (Seo et al., 2007; Shen and Phanikumar, 2010; Xu et al., 2012), is numerically more rigorous than the one-way coupling scheme. The main difference between the above two methods lies in the ability to conserve mass within a single step for back-and-forth information exchange. In an iterative method, the head/flux boundaries are iteratively exchanged. There is a cost-benefit tradeoff to obtain higher numerical efficiency.

During the late stages of the recharge in scenario 1 of case 1, the groundwater table rises quickly, which increases the burden on the coupling scheme. In our work, feedback iteration was conducted to eliminate the coupling error within the back-and-forth boundary exchange. To investigate how the feedback iteration influences the numerical accuracy as well as computational cost, solutions were compared with different closure criteria, instead of different maximal numbers of feedback iterations. For this purpose, scenario 1 in case 1 is tested with a range of closure criteria indicated by Closure = 0.001, 0.01, 0.1, 1, 5, and 20. Specifically, Closure = 20 (i.e., $\varepsilon_H = 20$ cm) is too large to regulate any feedback iteration, and is thus replaced by “non-iterative”. The ε_F , indicating the closure of the Neumann boundary feedback iteration, is usually related to the phreatic Darcian flux. To avoid its impact on the discussion below, we assume $\varepsilon_F = +\infty$, which means no regulation from the flux boundary

exchange. Due to less dynamic in the groundwater sub-model, the empirical relaxation factors were both set by 1.0 to have
365 straight forward update of the interfacial boundaries, i.e., z_t and F_{top} .

When the wetting front approached the phreatic surface (at $t = 2.4$ d), the number of feedback iteration increased dramatically, see **Figure 11a**. This was caused by the intensive rise of the water table within each macro time step ΔT . The head/flux interfacial boundaries were thus not easy to approximate the “truth”. With several attempts to exchange the head/flux boundaries, the head solution was effectively drawn towards the “truth”, see **Figure 11b**. With Closure < 2 , i.e., $\varepsilon_H < 2$ cm, the
370 coupling errors were significantly reduced, see **Figure 11c**. The cost-benefit curve, which was quantified by the number of feedback iteration and RMSE, was indicative to problems with larger scales, and higher dimensionalities.

4.4 Parsimonious decision making

The feedback coupling schemes, either iteratively or non-iteratively, increase the degree of freedom for the users to manage the sub-models with different governing equations, numerical algorithms, as well as the heterogeneities in parameters and
375 variabilities in hydrologic stresses. For practical purposes, a significant concern is how to efficiently handle the complicated and scale-disparate systems.

For problems with rapid changes in groundwater flows, as in case 2, the hydraulic gradient at the phreatic surface was large. Using a single soil column for such a complex situation introduced significant coupling errors at the water table, see **Figure 12a**. Although more sub-zones portioned means higher accuracy for the coupling method, five or more soil columns were
380 adequate enough approaching the “truth”. Furthermore, for the saturated nodes deep in the aquifer, such difference in coupling errors were of minor influence, see **Figure 12b**.

In case 3, a simple pumped and irrigated region was simulated with different numbers of soil columns. A range of tests with total numbers of 16, 12, 9, 5, and 3 soil columns were carried out to obtain a cost-benefit curve shown in **Figure 13c**. When partitioning the vadose zone into more than 12 soil columns, there was a slight reduction in solution errors (RMSE) and a
385 significant increase in computational cost caused by solving more 1D soil water models. Although parallelled computation could further reduce the numerical cost, representing the vadose zone with 3 sequentially calculated soil-water models achieved acceptable accuracy, as presented in **Figure 13a** and **b**. The computational cost for obtaining the fully-3D solution with VSF was 15.561 s, which was more than 11 times larger than an iterative feedback coupling method with soil-water models sequentially solved. Problems in more complicated real-world situations can thus be simplified to achieve higher numerical
390 efficiency.

4.5 Regional application

The Prudic et al.’s problem was originally designed to validate a streamflow routing package (Prudic et al., 2004). Stressed by

soil-surface infiltration, pumping wells, and general head boundary, the synthetic case was used to evaluate several unsaturated flow packages for MODFLOW (Twarakavi et al., 2008). Based on their studies, in case 4, we compared the developed iterative feedback coupling method with HPM. The hydraulic conductivity, as well as its heterogeneity, were forced to be consistent within the saturated and unsaturated zones, which is different from the case in Twarakavi et al (2008). **Figure 14a** gives the contours for the final phreatic head solutions, indicating a good match of the phreatic surface with the HYDRUS package. **Figure 14b-e** present the absolute head difference of the method developed here and the HYDRUS package at the end of stress periods 3, 6, 9, and 12. The dark color blocks indicated the largest difference in head solution. According to **Figure 6d**, the saturated grid cells controlled by the soil columns of #3、#9、#10、#15、#19 were suffering from the largest deviation, although with the same horizontal partitioning of the unsaturated zone. The strict iteratively two-way coupling contributes to such accuracy improvement.

For unsaturated-saturated flow situations, the vadose zone flow is important. **Figure 15** presents the water content profiles at sub-zones 1, 3, 5, 7, and 9 as examples. The solution obtained from the developed model matched well with the original HPM solution. For practical purpose, the manually controlled stress periods for the unsaturated sub-models are no longer constrained. In our method, the soil water models run at disparate numerical scales, which makes it possible to handle daily or hourly observed information rather than a stress period lasting 2 or more days in traditional coupled models.

5 Summary and conclusions

Fully-3D numerical models are available but are numerically expensive to simulate the regional unsaturated-saturated flow. The quasi-3D method presented here, in contrast, with horizontally disconnected adjacent unsaturated nodes, significantly reduces the dimensionality and complexity of the problem. Such simplification brings about computational cost-saving and flexibility for better manipulation of the sub-models. However, the non-linearity in the soil-water retention curve, as well as the variability in realistic boundary stresses of the vadose and saturated zones, usually result in the scale-mismatching problems when attempting numerical coupling. In this work, the soil-water and groundwater models were coupled with an iterative feedback (two-way) coupling scheme. Three concerns about the multi-scale water balance at the phreatic interface are addressed using a range of numerical cases in multiple dimensionalities. We conclude:

(1) A new HYDRUS package for MODFLOW was developed by switching the θ and h forms of Richards' equation (RE) at each numerical node. The *switching-RE* circumvents the disadvantages of the h - and θ -form REs to achieve higher numerical stability and computational efficiency. The one-dimensional *switching-RE* was employed to simulate the rapid infiltration into a dry-sandy soil, and the swiftly altering atmospheric upper boundaries in a layered soil column. Compared with the h -form RE , the *switching-RE* used 10^5 times larger minimal time-step size (Δt_{min}) and conserved mass better. Lowering the non-

linearity of soil-water models with this switching scheme was promising for coupling complex flow modeling systems at regional scale.

425 (2) Stringent multi-scale water balance analysis at the water table was conducted to handle scale-mismatching problems, and to smooth the information delivered back-and-forth across the interface. In our work, the errors originating from inadequate phreatic boundary predictions were reduced firstly by a linear extrapolation method, and then by an iterative feedback. Compared with the traditional stepwise extension method, the linear extrapolation significantly reduced the coupling errors caused by scale-mismatching. For problems with severe soil-water and groundwater interactions, the coupling errors were significantly reduced by using an iterative feedback coupling scheme. The multi-scale water balance analysis mathematically
430 maintained numerical stabilities in the sub-models at disparate scales.

(3) When a moving phreatic boundary was assigned to the soil columns during the phreatic water balance analysis, it avoided the coupling errors originating from the saturated lateral fluxes. In practical applications for regional problems, the fluxes into and out of the saturated parts of the soil columns differed, which added to the complexity and phreatic water balance error of the coupling scheme. With a moving Dirichlet lower boundary, the saturated regions of the soil-water models were removed.
435 The coupling error was significantly reduced for problems with major groundwater flow. Extra cost-saving was achieved by minimizing the matrix sizes of the soil-water models.

Future investigation will focus on regional solute transport modeling based on the developed coupling scheme. Surface flow models, as well as the crop models, which appears to be less non-linear than the sub-surface models, will be coupled in an object-oriented modeling system. The RS- and GIS-based data class can then be used to handle more complicated large-scale
440 problems.

Data/code availability: All the data used in this study can be requested by email to the corresponding author Yuanyuan Zha at zhayuan87@gmail.com.

Appendix A

A.1 The moving Dirichlet lower boundary

445 The bottom node of a soil column is adaptively located at the phreatic surface, which makes it an area-averaged moving Dirichlet boundary

$$z_i(T) = \int_{s \in \Pi} H(T) ds / \int_{s \in \Pi} ds \quad (\text{A1})$$

where $z_i(T)$ [L] is the elevation of the water table; Π is the control domain of a soil column; $H(T)$ [L] is potentiometric head solution, as well as the elevation of the phreatic surface, which is obtained by solving the groundwater model; s is the horizontal

450 area.

To simulate the multi-scale flow process within a macro time step $\Delta T^{j+1} = T^{j+1} - T^j$, the lower boundary head of a soil column is temporally predicted either by stepwise extension of $z_t(T^j)$ (Seo et al., 2007; Shen and Phanikumar, 2010; Xu et al., 2012) or by linear extrapolation from $z_t(T^{j+1})$ and $z_t(T^j)$. In **Figure 2a**, the stepwise extension method ($z'_t(T^j)$) potentially causes large deviation from the “truth”. In our study, the linear extrapolation is resorted to reduce the coupling errors and to accelerate the convergence of the feedback iteration. The small-scale lower boundary head at time t ($T^j < t \leq T^{j+1}$) is given by

$$z_t(t) = \frac{(t - T^{j-1}) \cdot z_t(T^j) - (t - T^j) \cdot z_t(T^{j-1})}{T^j - T^{j-1}} \quad (\text{A2})$$

A.2 The Neumann upper boundary

The moving Dirichlet boundary introduces the need for water balance of a moving balancing domain above the water table (see **Figure 2b**), which is bounded by a specific elevation above the phreatic surface, z_s [L], and the dynamically changing phreatic surface, $z_t(t)$ [L].

Assume that the activated top layer in a three-dimensional groundwater model is conceptualized into a phreatic aquifer, the governing equation for this layer is given by

$$\bar{S}_y \frac{\partial H}{\partial t} = \frac{\partial}{\partial x} \left(K \bar{M} \frac{\partial H}{\partial x} \right) + \frac{\partial}{\partial y} \left(K \bar{M} \frac{\partial H}{\partial y} \right) + F_{top} - F_{base} \quad (\text{A3})$$

where \bar{M} [L] is the thickness of the phreatic layer, which is numerically defined as the layer below the vadoze zone, $\bar{M} = z_s - z_0$; z_0 is the bottom elevation of the top phreatic layer, $z_0 \ll z_s$; F_{top} [LT^{-1}] is the groundwater recharge into the activated top layer of the phreatic aquifer, $F_{top} = (K \cdot \partial H / \partial z)_{z=z_s}$; F_{base} is the leakage into an underlying numerical layer, $F_{base} = (K \cdot \partial H / \partial z)_{z=z_0}$ (positive downward, so as F_{top}). The long-term regional-scale parameter indicating the water yield caused by fluctuation of the water table (Nachabe, 2002), \bar{S}_y , [-], is calculated by

$$\bar{S}_y = V_w / (A \cdot \Delta H) \quad (\text{A4})$$

where V_w [L^3] is the amount of water release caused by fluctuation of the phreatic surface (ΔH [L]); A [L^2] is the area of interest.

A.3 The relaxed iterative feedback coupling

The relaxed feedback iteration method (Funaro et al., 1988; Mehl and Hill., 2013) is used to improve the convergence of head/flux at the phreatic surface, see **Figure A.1**. The Dirichlet lower boundary head for the soil columns, z_t , as well as the Neumann upper boundary flux for the phreatic surface, F_{top} , are updated within each iterative step (Niter)

$$\begin{aligned} z_t^{updated} &= \lambda_h \cdot z_t^{new} + (1 - \lambda_h) \cdot z_t^{old} \\ F_{top}^{updated} &= \lambda_f \cdot F_{top}^{new} + (1 - \lambda_f) \cdot F_{top}^{old} \end{aligned} \quad (\text{A5})$$

where superscript *old* (or *new*) indicates the previous (or newly calculated) head/flux boundaries at the coupling interface; λ_h and λ_f are the empirical relaxation factors for head/flux boundaries respectively. Their values are suggested to be within (0, 1].

The iteration ends when agreements are reached at

$$\left| z_i^{\text{updated}} - z_i^{\text{old}} \right| \leq \varepsilon_H \quad \text{and} \quad \left| F_{\text{top}}^{\text{updated}} - F_{\text{top}}^{\text{old}} \right| \leq \varepsilon_F \quad (\text{A6})$$

480 where ε_H [L] and ε_F [LT⁻¹] are residuals for the feedback iteration of interfacial head and flux.

Author contribution: Jicai Zeng, Yuanyuan Zha and Jinzhong Yang developed the new package for soil water movement based on a switching Richards' equation; Jicai Zeng and Yuanyuan Zha developed the coupling methods for efficiently joining the sub-models. Four of the co-authors made non-negligible efforts preparing the manuscript.

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628 **Table 1** Soil parameters used in the test cases.

| # | Soil | θ_r (cm ³ cm ⁻³) | θ_s (cm ³ cm ⁻³) | α (1/cm) | n | k_s (cm/d) |
|---|------------------------|--|--|-----------------|-------|--------------|
| 1 | Sand | 0.093 | 0.301 | 0.0547 | 4.264 | 504 |
| 2 | Berino loamy fine sand | 0.029 | 0.366 | 0.028 | 2.239 | 541 |
| 3 | Glendale clay loam | 0.106 | 0.469 | 0.010 | 1.395 | 13.1 |
| 4 | Loam | 0.078 | 0.430 | 0.036 | 1.560 | 24.96 |
| 5 | Sandy loam | 0.065 | 0.410 | 0.075 | 1.890 | 106.1 |

629

630 **Table 2** The precipitation, evaporation, and pumping rates in 12 stress periods.

| Stress period | Precipitation (mm/d) | ET (mm/d) | Pumping rate (m ³ /d) |
|---------------|----------------------|-----------|----------------------------------|
| 1 | 0.21 | 1.32 | 4078 |
| 2 | 1.69 | 1.32 | 4078 |
| 3 | 2.11 | 1.32 | 2039 |
| 4 | 4.21 | 1.32 | 2039 |
| 5 | 1.05 | 1.32 | 6116 |
| 6 | 2.11 | 1.32 | 0 |
| 7 | 0.63 | 1.32 | 4078 |
| 8 | 1.05 | 1.32 | 0 |
| 9 | 0.63 | 1.32 | 2039 |
| 10 | 0.42 | 1.32 | 0 |
| 11 | 0.21 | 1.32 | 6116 |
| 12 | 0.21 | 1.32 | 0 |

631

632 **Figure Captions**

633 **Figure 1:** Schematic of the space- and time-splitting strategy for coupling models at two independent scales. For a groundwater
634 model, spatial discretization is expected to be large ($\Delta x = 10^0 \text{ m} - 10^3 \text{ m}$); while for soil water models, it occurs to be small (Δx
635 $= 10^{-3} \text{ m} - 10^0 \text{ m}$). Multiple levels of temporal discretization are common for regional problems. For groundwater model, the
636 stress periods (SP) and macro time step sizes (ΔT) appear by months and days ($10^0 \text{ d} - 10^1 \text{ d}$). For soil water models, the time
637 step sizes are about $10^{-5} \text{ d} - 10^0 \text{ d}$.

638 **Figure 2:** The Dirichlet-Neumann coupling of the soil-water and groundwater flow models at different scales. (a) Linear or
639 stepwise prediction of Dirichlet lower boundary for the soil water flow model. (b) Water balance analysis based on a balancing
640 domain with moving lower boundary. Blue dash line is the linearly extrapolated groundwater table as an alternative for
641 prediction of Dirichlet lower boundary. J (or j), T (or t), and ΔT (or dt) are the time level, time, and time-step size at coarse
642 (or fine) scale. At any of the transient state (t), the balancing domain is bounded by a user-specified top elevation (z_s), and the
643 moving phreatic surface (z_t). At a transient time t (or T^j), the total mass volume in the moving balancing domain is indicated
644 by $w(t)$ (or $w(T^j)$). The saturated lateral flux of the moving domain is indicated by $l(t)$, while the unsaturated lateral flux is
645 neglected as the assumption of quasi-3D models. The water flux into and out of the balancing domain is indicated by q_{top} and
646 q_{bot} .

647 **Figure 3:** Rapidly changing atmospheric upper boundary conditions for scenario 2, case 1.

648 **Figure 4:** Schematic of the cross-sectional for test case 2. Two pumping wells with screens of $z = 0-200 \text{ cm}$ are located at x
649 $= 2,500 \text{ cm}$ and $5,000 \text{ cm}$. The pumping rates per unit width at well #1 and #2 are respectively $2 \times 10^4 \text{ cm}^2/\text{d}$ and $1 \times 10^4 \text{ cm}^2/\text{d}$,
650 respectively.

651 **Figure 5:** Different number of sub-zones partitioned for the quasi-3D simulations in Case 3. The vadose zone is partitioned
652 into 16, 12, 9, 5, and 3 sub-zones.

653 **Figure 6:** Input of the synthetic regional problem including (a) land surface elevation, (b) initial head, (c) bedrock elevation
654 of the aquifer, and (d) the sub-zones and boundaries.

655 **Figure 7:** The time-step sizes through the simulation of (a) sudden infiltration into a dry-sandy soil column, and (b) rapidly
656 changing atmospheric upper boundary conditions with a layered soil column.

657 **Figure 8:** Comparison of soil moisture content at $z = 0 \text{ cm}$, 50 cm , and 200 cm for the layered soil column with rapidly
658 changing upper boundary conditions (Scenario 2, Case 1). Taking the HYDRUS1D solution as the “truth”, RMSEs of solution
659 of the developed model are provided at different soil depth.

660 **Figure 9:** Water table changing with time for different macro time step sizes ($\Delta T = 0.005 \text{ d}$, 0.05 d , 0.1 d , and 0.2 d), in scenario
661 1, case 1. The HYDRUS1D solution is taken as the “truth”. Compared with the stepwise extended method (Seo et al., 2007),
662 the coupling error is significantly reduced by a linear prediction.

663 **Figure 10:** Comparison of RMSE of (a) the phreatic surface and (b) the head solution (at $z = 0$) between the moving-boundary
664 and the stationary-boundary methods. Three different lengths of the stationary soil columns, $L = 1,000$ cm, 500 cm, and 300
665 cm, are considered.

666 **Figure 11:** (a) The number of feedback iterations and (b) phreatic surface solution changing with different closure criteria. The
667 legend “Closure = 0.001” means $\varepsilon_H = 0.001$ cm is used to regulate the feedback iteration. The HYDRUS1D solution is taken
668 as “truth”. Tested in scenario 1, case 1.

669 **Figure 12:** Comparison of (a) water table and (b) head solution (at $z = 0$) that are changing by the number of soil columns.
670 Solutions obtained with a moving-boundary method in case 2.

671 **Figure 13:** (a) Comparison of contours of the phreatic surface solution obtained with the fully-3D and quasi-3D methods; (b)
672 Comparison of the phreatic surface at A-A' cross-section; (c) computational cost and RMSE changing by the number of total
673 soil columns.

674 **Figure 14:** (a) Comparison of elevation of the water table calculated by the HYDRUS package for MODFLOW (Seo et al.,
675 2007) and the developed method ($t = 365$ d); (b) The absolute head difference of the phreatic head solution by the method
676 developed here and HYDRUS package at the end of stress periods 3, 6, 9, and 12. (Case 4).

677 **Figure 15:** Comparison of water content profiles obtained from the HYDRUS package for MODFLOW (Seo et al., 2007) and
678 the developed iterative feedback coupling method. Sub-zones 1, 3, 5, 7, and 9 are shown as an example. ($t = 365$ d in Case 4).

679 **Figure A.1:** Flowchart of the relaxed iterative feedback coupling scheme. The relaxation is conducted at the interfacial
680 Dirichlet/Neumann boundaries during the feedback iterations (except for the time T^j)