

Understanding the Effects of Revegetated Shrubs on Fluxes of Energy, Water and Gross Primary Productivity in a Desert Steppe Ecosystem Using STEMMUS-SCOPE Model

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Abstract. Revegetation is one of the most effective ways to combat desertification and soil erosion in semiarid and arid regions. However, the perturbation of revegetation on ecohydrological processes, particularly its effects on the interplay between hydrological processes and vegetation growth under water stress, requires further investigation. This study evaluated the effects of revegetation on the energy, water and carbon fluxes in a desert steppe in Yanchi County, Ningxia Province, Northwest China, by simulating two vegetated scenarios (shrubs-grassland ecosystem and grassland ecosystem) using STEMMUS-SCOPE model. The model was validated by field observations from May to September of 2016-2019. The evaluation of revegetation effects relied on comparing simulated fluxes between two vegetated scenarios in 2016 and 2019. In both scenarios, turbulent energy was dominated by latent heat flux, which was stronger in the shrubs-grassland ecosystem (+ 7 %). Higher leaf area index and root water uptake of C3 shrubs (*Caragana Intermedia*) resulted in increased carbon fixation (+ 83 %) and transpiration (+ 72 %) of the shrubs-grassland ecosystem compared to C3 grassland ecosystem. Accompanied by a marked increase in root water uptake (+ 123 %), revegetation intensified water consumption beyond the levels of received precipitation. These results highlight the critical importance of considering both energy and water budgets in water-limited ecosystems during ecological restoration, to avert soil water depletion.

1 Introduction

Global efforts in revegetation have been made to combat climate change and desertification. For example, satellite data reveals that the revegetation programs in China have contributed about 10.5 % of the increased global greening during 2000-2017 (Chen et al., 2019). This large-scale revegetation program ('Grain-to-Green') was initiated to improve the ecosystem service of degraded desert steppe in northern China since the 1990s (Liu et al., 2021). On the one hand, it is proven effective in controlling soil erosion and enhancing carbon sequestration (Liu et al., 2021; Zhang et al., 2018). On the other hand, the

conflict between the water deficit and the development of the shrub community has become an increasing concern, particularly in arid and semiarid lands where ecosystems are fragile and suffering intense water stress (D’Odorico et al., 2012; Tian et al., 2017; Huxman et al., 2005; Zhang et al., 2018). For example, the revegetation in China’s Loess Plateau has increased the net primary productivity and evapotranspiration, but the ecosystem is approaching sustainable water resource limits (Feng et al., 2016). Specifically, field studies reported that the revegetation leads to the depletion of soil moisture (Liu and Shao, 2015; Jia et al., 2017), formation of the dry soil layer (Fu et al., 2012; Jia et al., 2017; Jian et al., 2015; Gao et al., 2023) and reallocation of the energy partitioning along with changes in vegetation distribution and canopy structure (Chen et al., 2015). Therefore, implementing revegetation programs but ignoring their long-term effects on energy, water and carbon balance may act contrary to ecologically sustainable development. Quantitative assessment of energy, water, and carbon fluxes is essential for evaluating the impact of revegetation, including the determination of water resource limits and optimal plant coverage for revegetation (Fu et al., 2012; Feng et al., 2016). However, the lack of long-term observations makes it difficult to reproduce the energy, water and carbon cycles of the ecosystems before and after the revegetation practice. To overcome this challenge, process-based land surface models (LSMs) can provide a better understanding of the energy-water-carbon flows of ecosystems (Du et al., 2021; Gong et al., 2016).

The past few decades have seen the rapid development of LSMs for dryland ecosystems based on the soil-vegetation-atmosphere transfer continuum (SVAT) (Tague et al., 2004; Ivanov et al., 2008; Fatichi et al., 2016; Niu et al., 2020). It is widely believed that the dominant constraint of vegetation development in the semiarid region is soil water availability, which manifests itself in regulating photosynthesis, evapotranspiration and root distribution (Camargo and Kemanian, 2016; Fan et al., 2017). In this context, accurate soil water modelling in LSMs is critical for the overall model performance in predicting energy, water and carbon fluxes. However, some existing deficiencies in the soil water modelling include: (a) computing the soil water content with a simple “bucket” approximation (e.g., RHESSys and Biome-BGC), (b) defining maximum root water uptake capacity with empirical constants (e.g., CLM and tRIBS + VEGGIE) or by using a direct function based on soil water availability (Zeng et al., 1998; Tague et al., 2004; Zhang et al., 2013; Fisher et al., 2014; Newman et al., 2006). The “bucket” model may overlook the soil water movement through different layers due to its simple representation of the vertical soil profile and rooting structure (Romano et al., 2011; Du et al., 2021). Moreover, the decoupling between soil states and vegetation traits, or simplified process representation, such as constraining the vegetation growth using a constant water stress factor, might lead to biased simulation in aboveground fluxes (Bayat et al., 2019).

Given the critical role of water availability in semiarid ecosystems, we need a mechanistic insight into ecohydrological processes, particularly the water movement across various soil layers and its influence on photosynthesis. Here, we advance the understanding of ecohydrological processes in a semiarid desert steppe ecosystem by applying a process-based model STEMMUS-SCOPE. STEMMUS represents Simultaneous Transfer of Energy, Momentum, and Mass in Unsaturated Soil, while SCOPE is for Soil-Canopy Observation of Photosynthesis and Energy. The coupled STEMMUS-SCOPE model can

simulate the profile of dynamic root length density for estimating root water uptake (RWU) and the hydraulic resistance from soil, to root, stem, and to leaf (Wang et al., 2021b; Zeng et al., 2011a, 2009b). Thanks to the inclusion of plant hydraulic connection between soil and leaf, the coupled model realizes the influence of soil moisture variation on photosynthetic and stomatal behaviors, which facilitates the investigation of water stress effects on vegetation functioning.

To explore the intricate interactions between vegetation and changing land surface conditions, we specifically investigate and answer the following questions: (i) Can the mechanistic model STEMMUS-SCOPE reliably represent the fluxes of energy, water and gross primary productivity before and after revegetation? (ii) Which fluxes are most affected by revegetation, and how do they respond to water water availability? (iii) Will the revegetated shrubs lead to soil water depletion? Answering these questions, we presented the following work in this study. First, the contributions and leaf area index of shrubs and grasses were defined (Section 2.3.1 and 2.3.2) and they were further used to construct two vegetated scenarios (grassland ecosystem and shrubs-grassland ecosystem) (Section 2.3.3). Secondly, the sensitivity of ten critical parameters was analyzed using the Morris method (Section 2.4.1 and Section 3.1). Thirdly, STEMMUS-SCOPE was calibrated and validated with observations over May-July in 2018 and May-September in 2016, 2017 and 2019, respectively (Section 3.2). Finally, the simulated fluxes from the two scenarios were compared and their differences were analyzed (Section 3.3).

2 Materials and Methods

2.1 Study site

The study area (107°29'37" E, 37°49'46" N) is located in Yangzhaizi village of Yanchi County, Ningxia Province, which is a typical agricultural-pastoral ecotone with a mid-temperate semiarid continental climate (Fig. 1). The mean annual air temperature (1958-2017) was 8.34 °C. The mean annual precipitation is 296.99 mm, and about 80 % of the rain falls between June and September (Jia et al., 2018). Since the 1990s, the grazing-prohibiting policy and the revegetation program have been implemented in the study area to combat desertification.

The study site is a 30 m × 30 m fenced plot characterized by revegetated shrubs and natural grassland. The shrub strips were planted at an interval of 6-7 m, and the average distance between two neighbouring shrubs in each strip was less than 1 m (Fig. S2). Between and below the canopy of shrubs, sparse grasses grow and soil moisture sensor (SM150, DELTA-T, UK), soil temperature sensor (107-L, BetaTherm, DE) and heat flux plate (HFP01, Hukseflux, NL) were installed at 10 cm soil depth under the grasses. The EC flux tower has an open path CO₂/H₂O analyzer (LI-7500, LI-COR Inc., USA) and a 3D ultrasonic anemometer (Wind Master Pro, Gill, UK), a tipping bucket rain gauge (TE525 MM-L, Texas Electronics, USA), net radiometer (CNR-4, Kippen&zonen, NL), temperature and relative humidity probe (HMP45C, CSL USA), and is surrounded by *Caragana intermedia* perennial shrubs and native herbaceous plants (Fig. 1). The active growing season of the shrubs and grasses lasts from May to September. The predominant soil texture is characterized as aeolian sandy soil.

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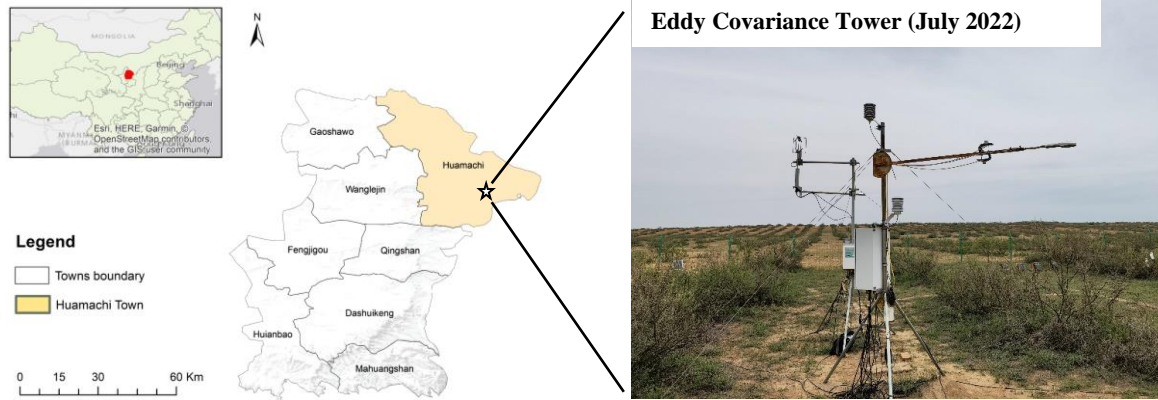


Figure 1. Overview of the study area. The red area in the map (from OpenStreetMap) is Yanchi County located in China. The white star denotes the field station located in Yangzhaizi village in Yanchi County, Ningxia Province.

2.2 STEMMUS-SCOPE model

105 Only the major modules in STEMMUS-SCOPE used in this study are described in Eqs (S1) - (S26). The original descriptions
of model components and other applications are elaborated by Bayat et al. (2019), Van der Tol et al. (2009), Wang et al.
(2021b), Yang et al. (2020), Zeng et al. (2011a, 2011b). The STEMMUS-SCOPE is employed to simulate the energy, water
and carbon fluxes for the two ecosystems, representing the scenarios before and after revegetation. The following assumptions
are made for modelling: (i) A land unit is structured as a vertical continuum, which consists of shrub or grass and soil column;
110 (ii) As the groundwater is more than 6 m below the surface in the study area (Du et al., 2021), the boundary condition at the
bottom of the soil column (i.e., 5 m depth) is set as gravity drainage; (iii) the soil texture is vertically homogeneous. In this
context, the model runs at a half-hourly time step, with the input of meteorological forcings, soil hydraulic and plant trait
parameters (Table S1). Model outputs are compared to data from the EC tower, which includes net radiation, latent heat flux,
sensible heat flux, and gross primary productivity. Additionally, ground measurements of soil moisture, soil temperature, and
115 corrected surface soil heat flux (using Eq. S9) are used for validation.

2.3 Simulation scenarios design

2.3.1 Land cover classification

Two vegetation covers both contribute to the EC observations and must be reflected in the model parametrization. Therefore,
120 a classification map is needed to derive the fractional vegetation cover for a more accurate modelling. The Supervised
Classification Method in ERDAS 2020 was used to determine the fractional cover of shrubs, grasses and bare soil based on an
image taken by unmanned aerial vehicle (Fig. S2).

STEMMUS-SCOPE considers the soil-root-canopy continuum, and quantifies the amount of energy received and water evaporated from its canopy and soil based on the leaf area index and leaf inclination (e.g., with four temperature variables: sunlit/shaded leaf temperatures, sunlit/shaded soil surface temperatures). Here, we assume that the 40 % coverage of bare soil distributed in each simulated soil-root-canopy continuum (i.e., either shrub or grass) and make the approximated contribution of 58.33 % for shrubland and 41.67 % for grassland (Table S3).

130 2.3.2 Reconstructed LAI

Leaf area index (LAI) is a critical variable in calculating the gross primary productivity (GPP) and latent heat flux (LE) in the STEMMUS-SCOPE. The MODIS 4-day LAI data (LAI_{MODIS}) during 2016-2019 was extracted from the Google Earth Engine Platform. Further, we applied the Harmonic Analysis of Time Series (HANTS) algorithm in MATLAB to smooth the outliers (Supplement Section 2).

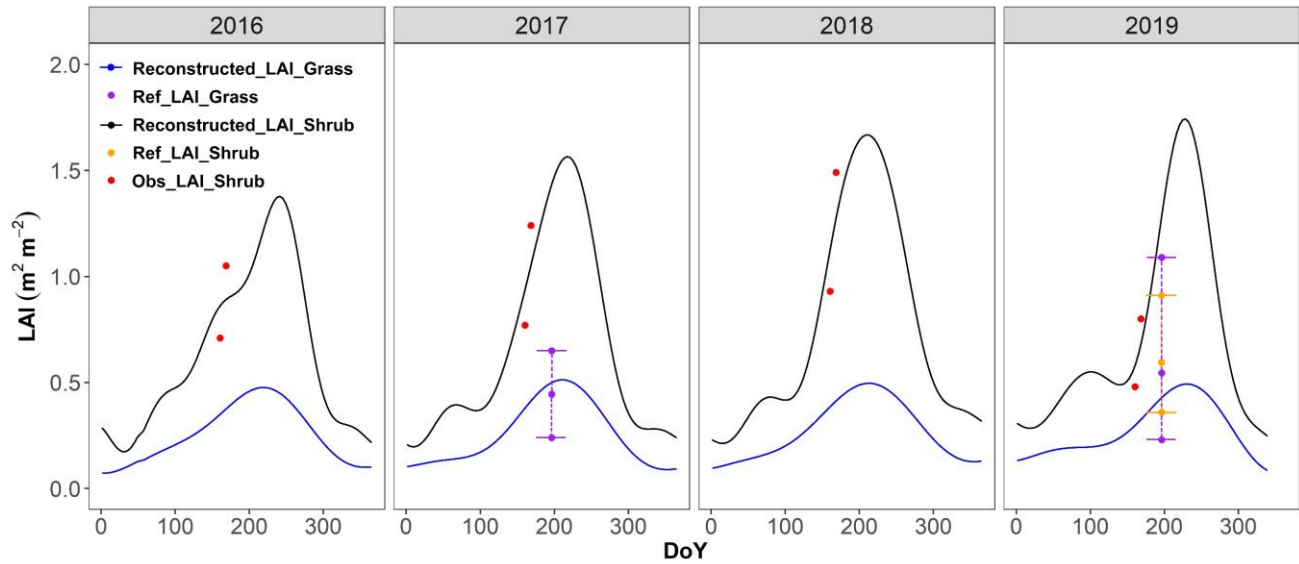
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The study site is a 30 m \times 30 m plot with two species, while LAI_{MODIS} can only provide an overall LAI with 500 m spatial resolution. To achieve the simulated flux partitioning from two land covers, we simulated shrub grid and grass grid with the LAI of shrub (LAI_{shrub}) and LAI of grassland (LAI_{grass}), respectively (Fig. 3c). With only two field measurements in 2022, LAI_{shrub} was corrected by multiplying smoothed LAI_{MODIS} by 2.33 (Table S5). LAI_{grass} was estimated by assuming it was $\frac{1}{4}$

140 of that of the shrubs, and by disaggregation of LAI_{MODIS} with the following conditions:

- i. $f_{shrub} * LAI_{shrub}(i) + f_{grass} * LAI_{grass}(i) + f_{baresoil} * LAI_{baresoil} = LAI_{MODIS}(i)$
- ii. $f_{shrub} + f_{grass} + f_{baresoil} = 1$
- iii. $LAI_{baresoil} = 0$
- iv. $LAI_{shrub}(i) \approx 4 LAI_{grass}(i)$ (Dan et al., 2020)
- 145 v. $LAI_{grass}(i)$ should follow the temporal pattern of $LAI_{MODIS}(i)$ and it was $\sim 0.5 \text{ m}^2 \text{ m}^{-2}$ (Yang et al., 2019; Dan, 2020).

where f_{shrub} , f_{grass} and $f_{baresoil}$ are the fractional cover of shrubs, grasses and bare soil, respectively. With the above constraints, LAI_{grass} shown on Fig. 2 was generated by the HANTS algorithm (Table S4 and Fig. S3).



150 **Figure 2. Reconstructed LAI of shrubland and grassland from 2016 to 2019. The red dots (Obs_LAI_Shrub) are the observed LAI of shrubs that were calculated based on the correlation derived from the observations in 2022 (Table S5). The purple dots and dotted lines (Ref_LAI_Grass) represent the ranges of reference LAI that were measured for the nearby grasslands (Yang et al., 2019; Dan, 2020). The yellow dots and dotted lines (Ref_LAI_Shrub) represent the ranges of reference LAI that were measured for the nearby shrublands (Dan, 2020).**

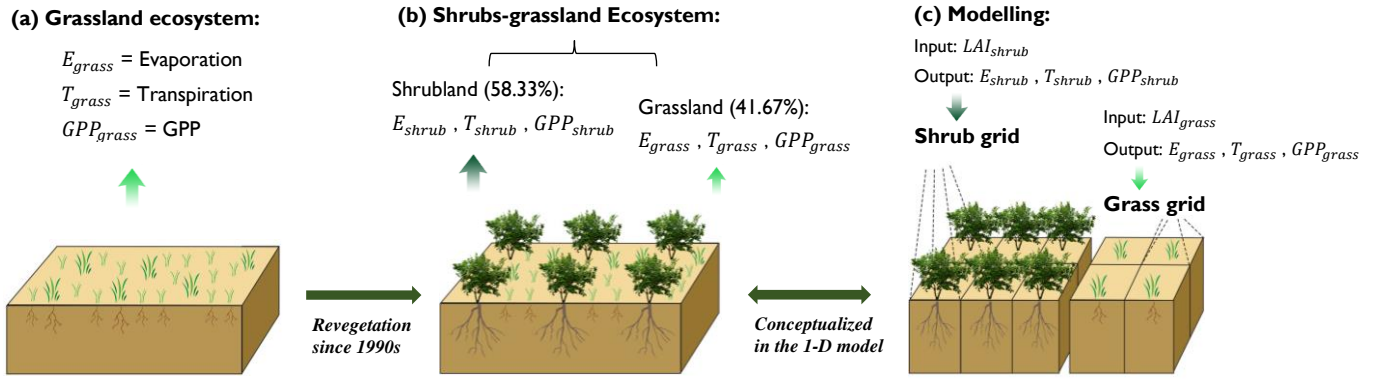
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2.3.3 Scenarios design

Two sets of canopy parametrization schemes (Table S1) together with the reconstructed LAI were used to simulate the fluxes of shrub grid and grass grid, respectively (Fig. 3c). We assumed the shrubland and grassland had the same meteorological environment; likewise, the same initial conditions of soil temperature and soil water content were assigned to the two land covers (Table S2). To accurately depict the fluxes of mixed surfaces containing both shrubs and grasses, the simulated fluxes from the shrubland and grassland simulations were partitioned based on their respective contributions, as demonstrated in Fig. 3(b). For instance, the total evaporation, transpiration and gross primary productivity (GPP) were the sum weighted by their contributions. As an example:

$$GPP = C_{shrub} GPP_{shrub} + (1 - C_{shrub}) GPP_{grass} \quad (1)$$

165 C_{shrub} is the contribution of shrubland to the overall flux, and its value is 58.33 % while the contribution of grassland is 41.67 % (Table S3). The same partitioning method as GPP was applied to net radiation, latent heat flux and sensible heat flux, in which the latent heat was converted into evaporation flux using the evaporation heat at 20 °C.



170 **Figure 3. Conceptual diagram of (a) native grassland scenario before revegetation (hereafter is denoted as grassland ecosystem), (b) realistic scenario with mixed shrubs and grasses (hereafter is denoted as shrubs-grassland ecosystem) and (c) conceptualization in the model. The bare soil evaporation is implicitly included in E_{shrub} and E_{grass} , where E is the evaporation.**

2.4 Model calibration and validation

2.4.1 Sensitivity analysis

175 Morris global sensitivity analysis was applied to evaluate the sensitivity of model simulations to the variation of input parameters. Morris SA, a screening-based method, can give a qualitative ranking of parameters at a relatively low computational cost (Herman et al., 2013). The most influential parameters for STEMMUS model and SCOPE model identified by other literature are: Maximum carboxylation rate (V_{cmax}), Ball-Berry stomatal conductance parameter (m), Leaf inclination (LIDFa), Residual soil water content (θ_r), Van Genuchten parameters (n and α), Saturated hydraulic conductivity (K_{sat}),
 180 Maximum rooting depth (Rdepth), Fitted extinction coefficient (β) and Root density (R_D) (Wang et al., 2021a; Verrelst et al., 2015; Jackson et al., 1997). Note that sensitivity analysis was only done for parameters of the shrubland simulation and the range of soil parameters was set for the specific sandy soil (Table 1).

The Morris analysis was achieved using the SALib package in Python and its workflow is explained in Supplement Section 3
 185 (Herman and Usher, 2017). Besides, 220 set of parameter trails generated 220 set of fluxes, which were compared with the observed fluxes from May to July in 2018. The performance metrics, including coefficient of determination (R^2), root mean square error (RMSE) and an objective function, were calculated for each trail. The objective function is calculated as

$$RMSE_{normalized} = \frac{RMSE_{SWC}}{\overline{Obs}_{SWC}} + \frac{RMSE_{LE}}{\overline{Obs}_{LE}} + \frac{RMSE_{GPP}}{\overline{Obs}_{GPP}}$$
, where the $\overline{Obs}_{SWC/LE/GPP}$ is the average values of observed SWC, LE and GPP throughout the investigation period, respectively (Groenendijk et al., 2011). The optimal trail for shrubland is identified
 190 by the lowest $RMSE_{normalized}$ and the highest R^2 .

Table 1. Range of critical parameters for sensitivity analysis.

Module	Parameters	Description	Units	Range
Canopy	V_{cmax}	Maximum carboxylation rate of C3 Shrub	$\mu\text{mol m}^{-2} \text{s}^{-1}$	[60 ^a , 250]
	m	Default values of Ball-Berry slope	-	[2, 20]
	LIDFa	Default values of leaf inclination	-	[-1, 1]
Soil	θ_r	Soil parameters of sandy soil in similar sites (e.g., with similar species and/or soil texture) in Northwest China	$\text{m}^3 \text{m}^{-3}$	[0.004 ^b , 0.035 ^c]
	n		-	[1.38 ^b , 2.09 ^b]
	α		m^{-1}	[0.0028 ^b , 0.03 ^d]
	K_{sat}		cm d^{-1}	[100, 300 ^b]
Root	Rdepth	Maximum rooting depth	cm	[67.5 ^b , 500]
	β	Fitted extinction coefficient	-	[0.909, 0.982] ^e
	R_D	Root density	g m^{-3}	[0.21 ^e , 0.492]

^a(Wang et al., 2017); ^b(Gong et al., 2016); ^c(Montzka et al., 2017); ^d(Wei et al., 2019); ^e(Jackson et al., 1997).

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2.4.2 Performance metrics

RMSE and R^2 were used to evaluate the quality of the model predictions.

$$RMSE = \sqrt{\frac{\sum_{i=1}^n (x_o - x_s)^2}{n}} \quad (3)$$

$$R^2 = 1 - \frac{\sum (x_o - x_s)^2}{\sum (x_o - \bar{x}_o)^2} \quad (4)$$

200 where x_s is the simulated value, x_o is the corresponding observed value and \bar{x}_o is the mean of observed values, n is the number of data records.

3 Results

3.1 Model sensitivity

Soil hydraulic parameters exert the strongest influence on simulated fluxes, with key parameters identified for each (Fig. S5):
 205 (i) LIDFa for net radiation (Rn), (ii) α and n for ground heat flux (G), (iii) m for latent heat flux (LE) and sensible heat flux (H), (iv) V_{cmax} for GPP, (v) α and n for soil water content (SWC). Maximum rooting depth (Rdepth) and root density (R_D) also have notable impacts on SWC and G simulations. Consistent with the STEMMUS-SCOPE model calculations, Rdepth and R_D primarily govern root distribution, thereby affecting root water uptake across various soil layers and ultimately impacting SWC. The parameter trail achieving the lowest $RMSEn$ includes V_{cmax} at 123, m at 6.8, LIDFa at -0.33 , α at
 210 0.005, n at 1.71, θ_r at 0.014 $\text{m}^3 \text{m}^{-3}$ and K_{sat} at 100 cm d^{-1} . The optimal trail was adapted for grassland scenario with adjustments to root parameters as detailed in Table S1.

3.2 Model performance

Data collected from May to July 2018 was used to calibrate the model (Fig. S6-S8). Compared with the simulations considering only one land cover, the model performance was better in capturing the dynamic and magnitude of energy and carbon fluxes when considering the mixed land covers (Fig. S8).

Data collected during May-September in 2016 and 2019, May-July in 2017 was used to validate the model simulations. The simulated energy fluxes showed satisfactory agreement with the observed values, with the R^2 all above 0.67 and with the RMSE ranging from 19.64 to 66.90 $W m^{-2}$ (Fig. 4a-4d). The simulated turbulent flux (i.e., LE+H) followed well the trend of the measured values ($R^2 = 0.86$, RMSE = 59.16 $W m^{-2}$).

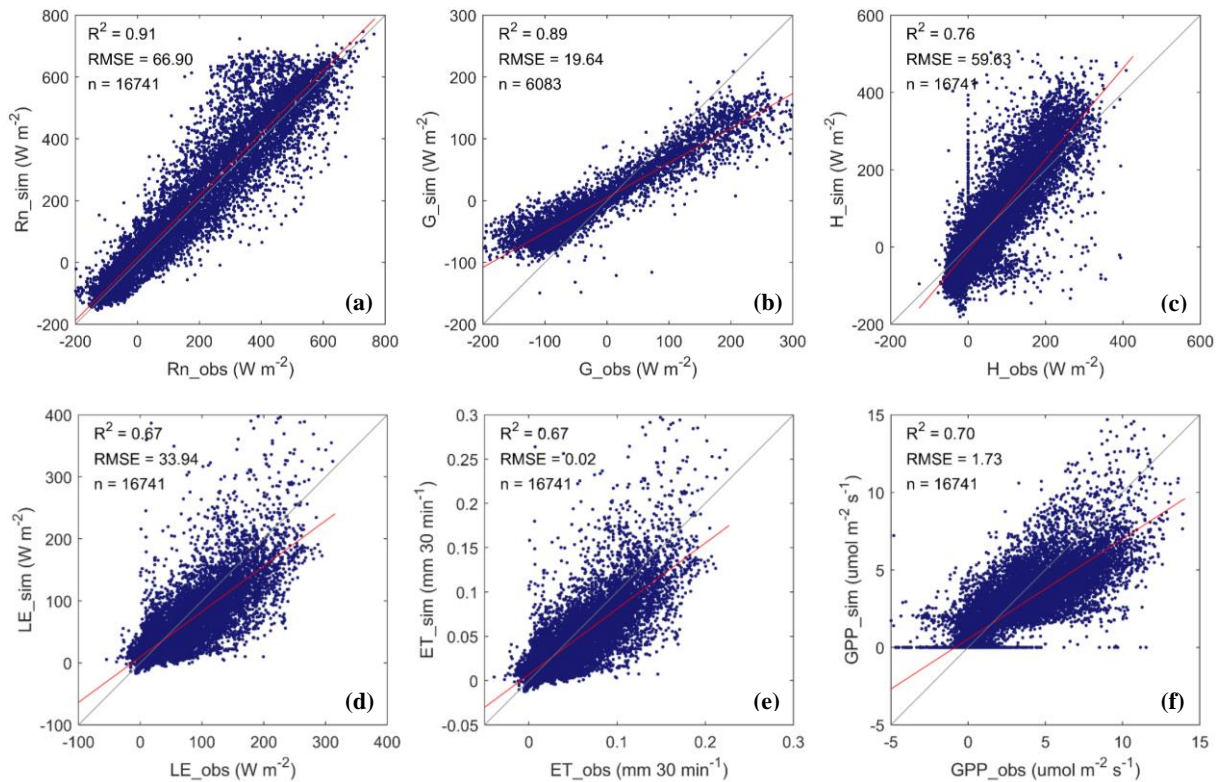


Figure 4. Comparing the observed (obs) and simulated (sim) half-hour values of (a) Net radiation (Rn), (b) Ground heat flux (G, where only 6083 data points in year 2019 was the valid measurements for comparison), (c) Sensible heat flux (H), (d) Latent heat flux (LE), (e) Evapotranspiration (ET) and (f) Gross primary productivity (GPP). The performance statistics are summarized in Table S6.

The simulated soil water content (SWC) and soil temperature (Ts) were primarily validated by the observed data from the grassland ecosystem (Fig. 5) as the sensors were installed under the grassland. The model can capture the SWC dynamics in response to each rainfall event (Fig. 5a: $R^2 = 0.84$; RMSE = 0.01 $m^3 m^{-3}$). In addition, the diurnal patterns were also captured

230 though their amplitudes were not as significant as observations. The simulated Ts at 10 cm depth also displayed an apparent diurnal pattern and was comparable with the observed Ts (Fig. 5b: $R^2 = 0.85$; RSME = 2.74 °C).

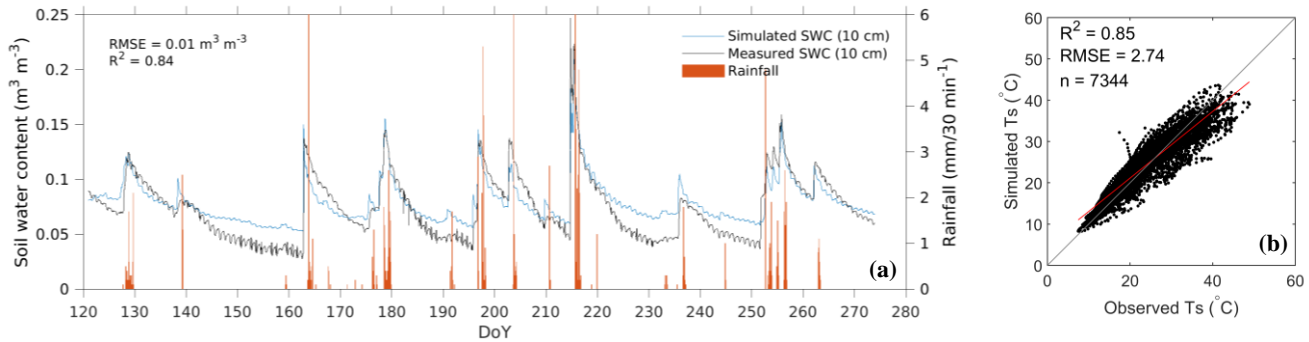


Figure 5. Model simulated and measured half-hourly soil water content (SWC) and (b) soil temperature (Ts) during May-September in 2019; (a) Temporal dynamics of SWC; (b) Scatter plots between the simulated and the observed Ts.

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3.3 Comparison between two scenarios

The averaged values of simulated fluxes during May-September in 2016 and 2019 were used to understand the diurnal and daily variations of energy fluxes. Flux differences between the shrub-grassland and grassland ecosystems were determined by subtraction, with mean values and seasonal totals presented as mean (\pm standard deviations) in Tables S7-S9.

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3.3.1 Diurnal variations of energy fluxes

The envelope of net radiation (R_n) ranged from -91.61 W m^{-2} to 505.69 W m^{-2} (Fig. S9 (a)). The shrubs-grassland ecosystem was likely to receive more radiance because of the denser canopies of shrubs, reflected by the larger LAI in the model. However, differences in diurnal R_n between the two ecosystems were very small and mainly occurred at midday (i.e., 10:00 - 15:00 h), with an averaged difference of $23.23 \pm 4.33 \text{ W m}^{-2}$. During nighttime hours (17:00 - 07:00 h), the R_n was nearly identical for two ecosystems, showing negative values ($\sim -40 \text{ W m}^{-2}$) due to outgoing longwave radiation from soil and leaves.

The sensible heat flux (H) followed a similar pattern like the R_n but with a larger difference between the two ecosystems. The H reached the peak at ~ 262 (200) W m^{-2} in May of the shrubs-grassland (grassland) ecosystem (Fig. S9 (b)). During nighttime, H was below zero, indicating a heat transfer from the atmosphere to the ground due to the lower surface temperature of soil and canopy. The H of shrubs-grassland ecosystem appeared to be larger than that of grassland ecosystem whereas the seasonal H/R_n partitioning was similar ($\sim 37\%$) for both ecosystems. Latent heat flux (LE) had the maximum diurnal peak at 156 W m^{-2} in August of both ecosystems (Fig. S9 (c)). The LE steadily increased from June to August and dropped in September, which was in line with the plants growing stages and rainfall pattern.

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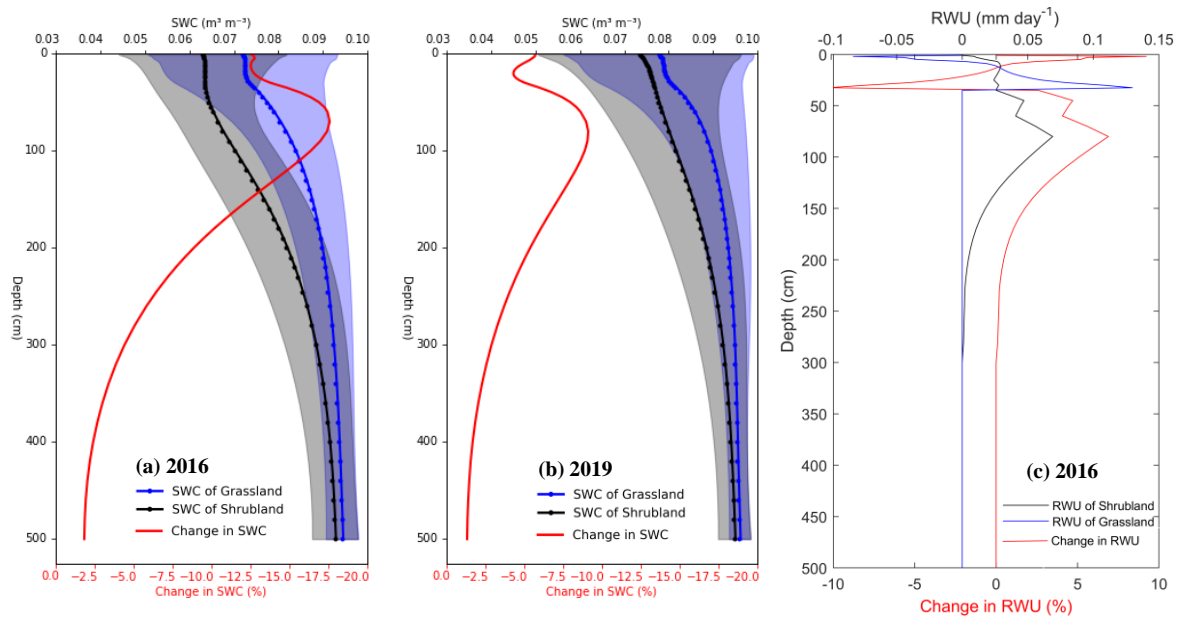
During May-September, the ground heat flux (G) peaked at ~ 11:00 am, with the averaged values of $133.10 \pm 6.03 \text{ W m}^{-2}$ ($161.57 \pm 7.72 \text{ W m}^{-2}$) for the shrubs-grassland (grassland) ecosystem (Fig. S9 (d)). More heat is transported through the surface and soil in the grassland ecosystem because less vegetation coverage induces more energy exchange in the soil. Throughout the growing seasons, more energy was stored in the soil under the grassland ecosystem (13.51 W m^{-2}) than that under the shrubs-grassland ecosystem (9.85 W m^{-2}).

3.3.2 Daily variations of water fluxes

We compared daily variations of soil water content (SWC), evaporation and transpiration between the grassland ecosystem and shrubs-grassland ecosystem, respectively. Additionally, we assessed the impact of rainfall on these water fluxes by comparing 2016 and 2019. The study categorized 2016 (seasonal rainfall of 218.1 mm) as a *dry* year and 2019 (292.4 mm) as a *normal* year, using the mean annual precipitation (296.99 mm) as a baseline.

Water fluxes: SWC and RWU

Generally, the SWC decreased in every soil layer after planting shrubs (Fig. 6a and 6b). Significant reductions in SWC (18% and 9% in 2016 and 2019, respectively) occurred between 50 cm depth to 100 cm depth, where most of the fine roots of shrubs concentrate (Jia et al., 2012; Zhu and Wang, 2020). Out of the root zone (i.e., below the 269 cm depth), the changes in SWC became less variable. Likely, the root water uptake of grasses (shrubs) increased from the surface layer and then decreased to zero at 35 cm (280 cm) depth. Moreover, the model successfully captured hydraulic redistribution, as indicated by negative RWU values in the relatively shallow root zone (Kennedy et al., 2019; Wang et al., 2021b). The negative RWU values resulted from the higher root water potential (in absolute value) compared to the soil water potential when the surface was too dry in the study area. When comparing the changes in RWU, the substitution of shrubs reduced RWU at the 13-33 cm depth while increased it at the 1-11 cm and 35-280 cm depth (Fig. 6c).



280 **Figure 6. Profile of the changes in soil water content (SWC) and root water uptake (RWU) after planting shrubs, which is calculated by using averaged value under the shrubland (black line) minus averaged values under the grassland (blue line) over the May to September in (a) 2016, (b) 2019 and (c) RWU for 2016 (RWU for 2019 has similar pattern). Note that the comparison was carried out between grassland ecosystem and shrubland ecosystem, instead of shrubs-grassland ecosystem.**

285 The SWC at 10 cm soil depth (hereafter denoted as surface SWC) under grassland peaked at $\sim 0.15 \text{ m}^3 \text{ m}^{-3}$ in July and August, followed by the most frequent rainfall events during the year (Fig. 7a). The surface SWC under both land covers showed rapid responses to rainfall, with sharp increases observed within just a few hours (Fig. 7a). At 100 cm soil depth (Fig. 7b), the SWC was less influenced by water exchange between the soil and atmosphere (i.e., evaporation and precipitation) (Zeng et al., 2009a; Yang and Fu, 2017; Zeng et al., 2009b).

290 The SWC under the shrubland was generally lower than under the grassland. Moreover, the average SWC difference at 100 cm depth was more pronounced, at $0.014 (0.008) \text{ m}^3 \text{ m}^{-3}$, than at the surface SWC with the values of $0.009 (0.004) \text{ m}^3 \text{ m}^{-3}$ in 2016 (2019) (Table S7). These contrasts indicate that the revegetation decreased the SWC and had a more significant effect on the SWC at deep soil layers. Particularly, in the dry year 2016, the differences in SWC between two land covers were more prominent and the shrubs drawn more soil water than that in 2019 (Fig. 7: left against right). This reflects that shrubs can tap
 295 into deeper soil water reserves through their developed root systems when experiencing drought (Wang et al., 2018; Zhu and Wang, 2020; Zhang et al., 2020b).

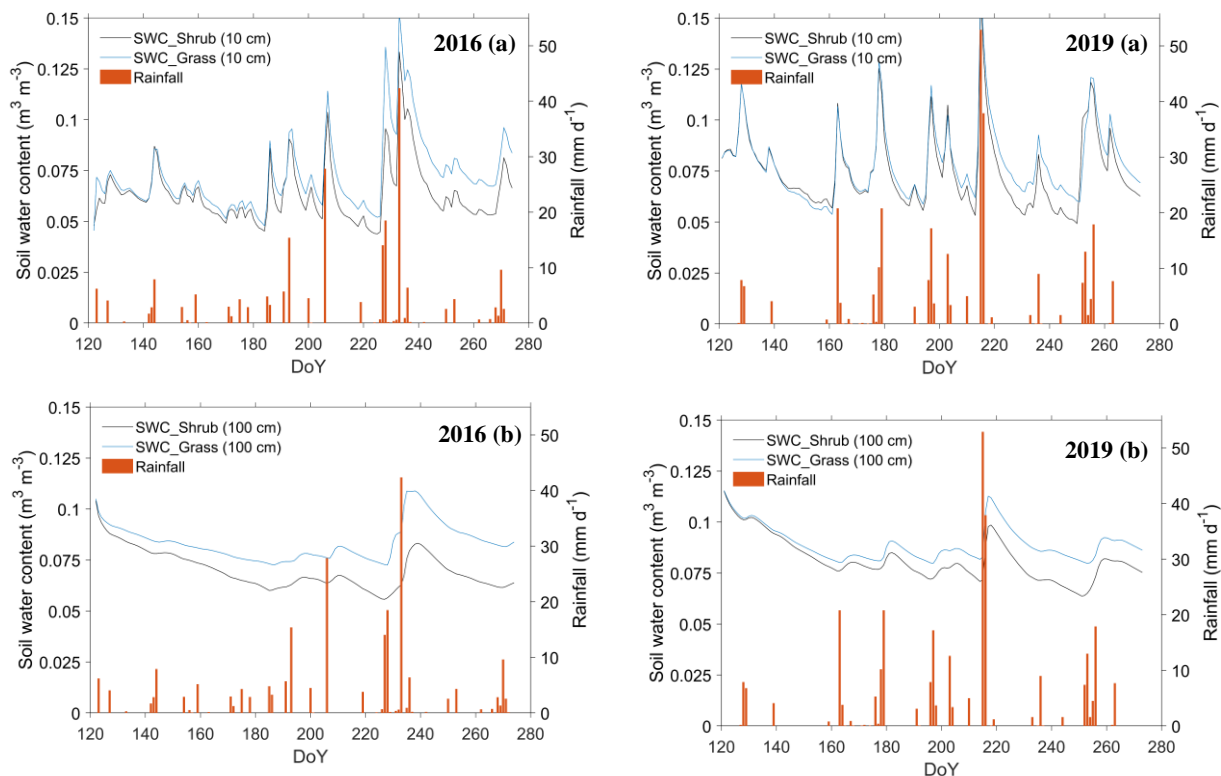


Figure 7. Daily variation of simulated SWC at (a) 10 cm and (b) 100 cm depth under two land covers during May–September in year 2016 (left) and 2019 (right). In the legend, ‘shrub’ denotes the shrub grid modelling.

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Water fluxes: Evaporation, Transpiration and ET

As shown in Fig. 8, the evaporation and transpiration boosted along with the rainfall pulses and their large day to day variation was noted. The seasonal evaporation of the shrubs-grassland ecosystem was reduced by 21 % (15 %) compared to that of the grassland ecosystem over the growing season in 2016 (2019) (Table S8). In contrast, the seasonal transpiration of the shrubs-grassland ecosystem was increased by 69 % (75 %) compared to that of the grassland ecosystem in 2016 (2019). Over the growing season in 2016 (2019), transpiration accounted for 48 % (41 %) of ET from the shrubs-grassland ecosystem, compared to 30% (25%) from grassland ecosystem. The seasonal ET of the shrubs-grassland ecosystem was 6 % (8 %) higher than that of the grassland ecosystem in 2016 (2019). In conclusion, the revegetated shrub reduced the evaporation but increased the transpiration, leading to a net increase in total ET at the ecosystem scale.

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As rainfall increased by 74.3 mm in 2019, the seasonal ET increased by 59.50 mm (20.88 mm) of the shrubs-grassland (grassland) ecosystem. However, the contributions of evaporation (~ 73 %) and transpiration (~ 27 %) to total ET remained stable for grassland ecosystem in two years. Besides, a slight variation was observed in SWC at 10 cm and 100 cm soil depth

under grassland between two years (Table S7). It seems possible that the water consumption of grassland was relatively stable
315 regardless of changes in precipitation amount (see discussion in [Section 4.3](#)).

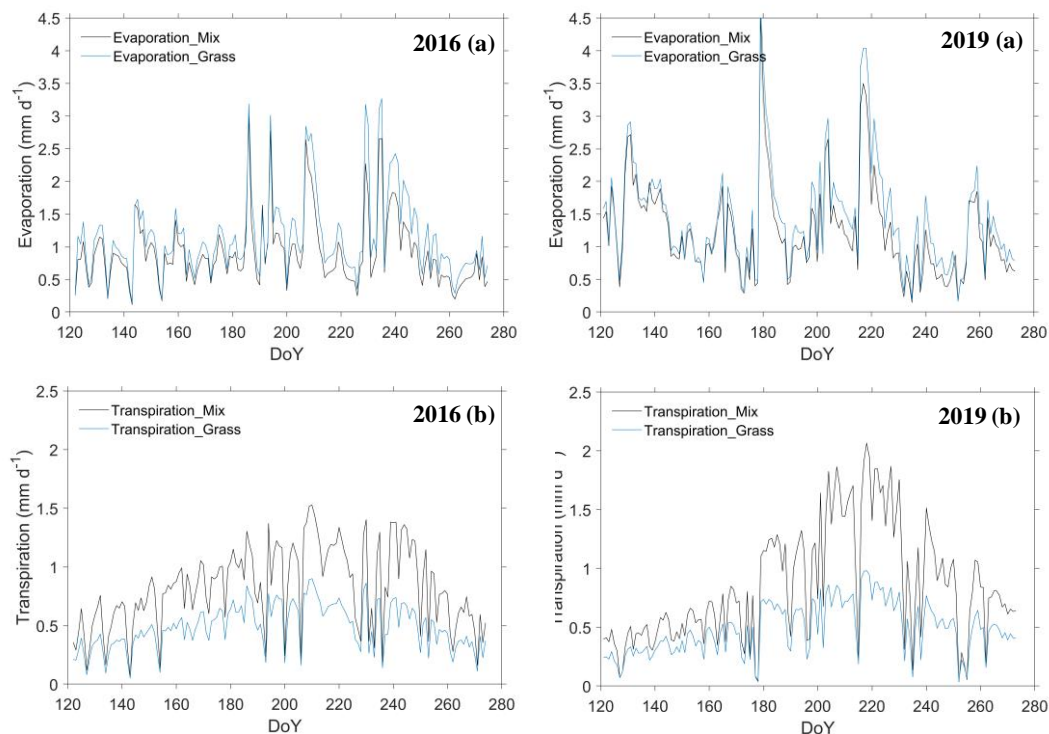


Figure 8. Daily variation of simulated (a) Evaporation and (b) Transpiration of two ecosystems during May–September in year 2016 (left) and 2019 (right).

3.3.3 Diurnal and daily variations GPP

320 The Gross Primary Productivity (GPP) of both ecosystems displayed evident diurnal patterns (Fig. S10 (a)). During the daytime, GPP was positive, indicating that the plants were taking up carbon. The plants halted their photosynthesis at nighttime with zero value of GPP. The magnitude of the diurnal variations was the largest in August, during which the GPP peaked at $7.36 \mu\text{mol m}^{-2} \text{s}^{-1}$ and $3.63 \mu\text{mol m}^{-2} \text{s}^{-1}$ during midday (10:00-14:00 h) of shrubs-grassland and grassland ecosystems, respectively. The GPP was consistently higher in the shrub-grassland ecosystem, particularly in July and August.

325 Daily GPP of the shrubs-grassland ecosystem tended to be more sensitive to the rainfall with more significant fluctuations, represented as the apparent increase in the DOY 207, 229 and 234 in year 2016 and DOY 180, 199, 217 in year 2019 (Fig. 9). Moreover, the shrubs-grassland ecosystem assimilated 76 % (89 %) more carbon than the grassland ecosystem over the growing season in 2016 (2019) (Table S9). In 2019, the seasonal GPP of shrubs-grassland ecosystem (grassland ecosystem)

increased by 15 % (7 %) compared to that in 2016. In the presence of more rainfall, the shrubs suffered less water stress,
 330 directly promoting the carboxylation rate for greater photosynthesis (Eq. S18).

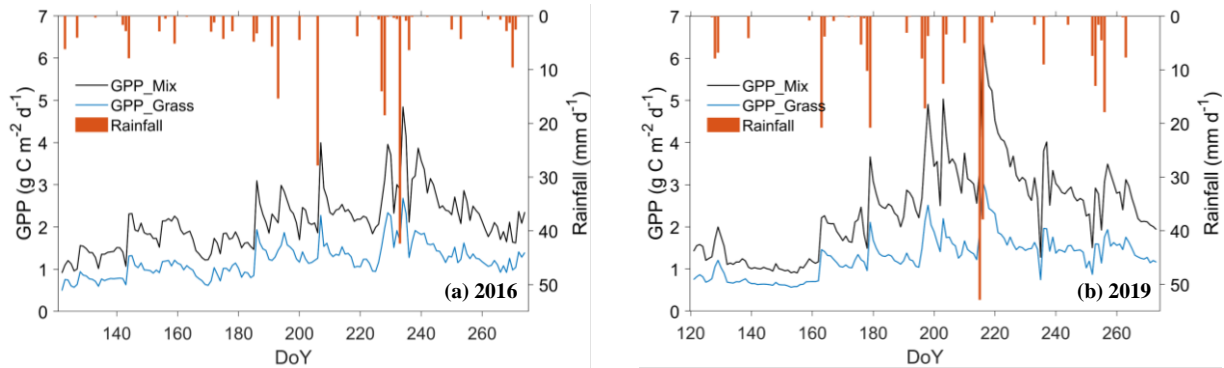
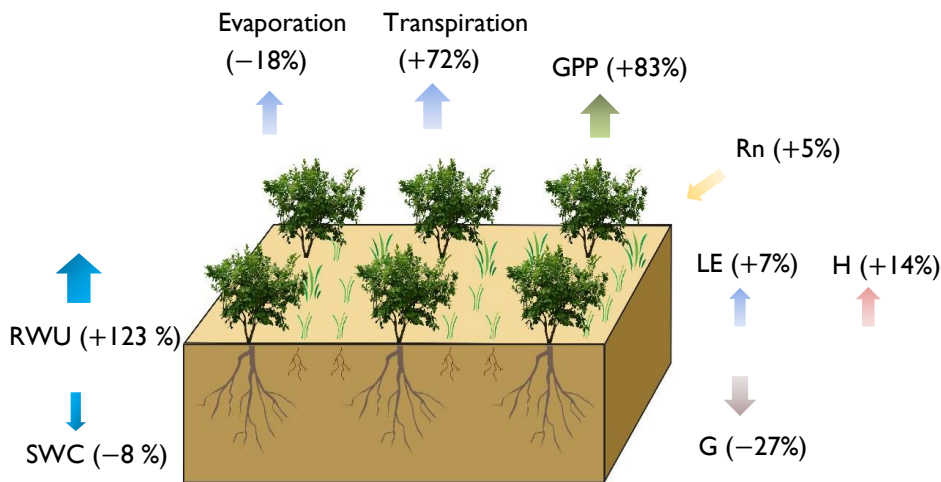


Figure 9. Daily variation of simulated Gross Primary Productivity (GPP) of two ecosystems during May–September in (a) 2016 and (b) 2019.

3.3.4 Changes caused by revegetation

335 Overall, our results show that revegetation increased R_n , LE and H by 5 %, 7 % and 14 %, respectively, while decreasing G by 27 %. The soil water content in the soil column was reduced by 8% and the root water uptake was increased sharply by 123%. The revegetation decreased the evaporation by 18%; meanwhile, the transpiration increased by 72 %. As for carbon flux, the revegetated shrubs increased the GPP by 83 % (Fig. 10).



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Figure 10. The changes in the energy, water and carbon fluxes caused by the revegetation, illustrated by the difference in averaged fluxes between shrubs-grassland ecosystem and grassland ecosystem. The arrows indicate the direction of the fluxes and the symbol ‘-’ represents a decrease while the ‘+’ represents an increase based on the grassland scenario.

4 Discussion

345 4.1 Evaluation of model performance and uncertainties

As summarized in Table S6, the model simulations showed a good match with the observed fluxes, but still, there are some deviations. Firstly, these deviations primarily stem from the model inputs and the scenarios design. For instance, (i) the estimated parameters (e.g., rooting depth and V_{cmax}) and reconstructed LAI for shrubland and grassland were critical for LE and GPP modelling; (ii) concerning the scenarios design, the contributions from shrubland and grassland that estimated from
350 only one UAV photo might not be representative over the investigation period because vegetation coverage is highly dependent on precipitation supply in the study area. The contributions from shrubland and grassland implicitly included the fraction of bare soil. The model then simulates evaporation from the soil beneath sunlit or shaded leaves and upscales it into a canopy/grid by multiplying LAI, rather than directly calculating evaporation from the bare soil. However, this approach may lead to an underestimation of soil evaporation, as a separate simulation of bare soil was not conducted in this study. To overcome this,
355 the model will be further developed to allow parallel computations of bare soil, low and high vegetations; (iii) direct comparison between simulated SWC and observed SWC under grassland leads to the bias because the observed SWC is the average of shrub, grass, and bare soil. However, SWC is a state variable that is not reasonable to aggregate/average from shrub, grasses and bare soil simulation in the current modelling scheme. Nonetheless, the comparison between simulated SWC and the observed value for either shrubland ($R^2 = 0.79$ and 0.82 in 2018 and 2019, respectively) or grassland ($R^2 = 0.88$ and 0.84
360 in 2018 and 2019, respectively) proves the credibility of model simulations; (iv) the quality of gap-filled forcing data and in-situ measurements are the basis for a valid comparison between simulations and observations. Although we applied a pre-assessment and filtering strategy in the energy fluxes with available data (Eq. S10), the energy closure issue results in uncertainty in comparing the H and LE/ET values, especially in years 2016 and 2017 without soil heat flux for quality control on energy fluxes.

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For the model itself, it is a one-dimensional vertical model without considering the lateral flow, which is assumed reasonably to be negligible in this flat study area. Nevertheless, at another site in Yanchi County with similar land covers, Gong et al. (2016) found that the absence of horizontal exchanges of water, vapour and heat advection in the model led to an underestimation of energy fluxes. However, there is always a trade-off between model complexity and effectiveness. And for
370 our focus on understanding the impact of revegetation on energy, water and carbon fluxes, the neglect of lateral flow in the STEMMUS-SCOPE model will not affect our conclusions.

Validation performances of shrubs-grassland ecosystem LE and GPP ($R^2 = 0.67$ and 0.70) were reasonable. Most of the discrepancies are represented as the overestimate of LE during large and continuous rainfall events. For instance, in 2016, the
375 model generated outliers during DOY 228 - 240 along with 69.56 mm rainfall within this period; likewise, the same was also reflected in 2017 and 2019 (Fig. S11). The overestimation of energy-limited evaporation rate, driven by the skin temperature,

is the main reason for the overestimation of LE. The causes of the outliers might be attributed to two factors: First, the use of a fixed time step (i.e., 30 minutes) in running STEMMUS-SCOPE was coarse temporally to achieve a precise numerical solution. Alternatively, the measurements of LE during rainfall events are less reliable.

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During the dry period without rainfall, the model tended to underestimate LE and GPP. As shown in Fig. S12, GPP simulations were notably low during water-stressed periods, where the water stress factor (WSF) approached zero. Several factors partially contribute to the underestimations in LE and GPP. Above ground, GPP is positively influenced by WSF, which in turn regulates V_{cmax} (Eq. S18). This affects both stomatal conductance and LE, making errors from approximated LAI (Section 2.3.2) and V_{cmax} potentially significant. Below ground, observations from other studies suggest that shrubs could adapt their RWU strategies, tapping into deep soil or using lateral roots. Unfortunately, the STEMMUS-SCOPE model lacks the flexibility to simulate these dynamic RWU strategies; it relies on initial SWC profile and root parameters to simulate root growth and RWU. In the STEMMUS-SCOPE model, the initial SWC profile not only determines the pattern of soil water storage in the soil column but also indicates the pattern of RWU. However, we lack the observations of SWC in deeper layers and root distributions. To mitigate the uncertainties caused by critical parameters, we carried out a sensitivity analysis to identify their influences and provide reference values. Our findings underline the importance of accurate LAI, V_{cmax} , Ball-Berry stomatal conductance parameter and multi-layer soil moisture for improving LE and GPP simulations.

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4.2 Effects of anthropogenic revegetation on ecosystem processes

395 4.2.1 On energy fluxes

The difference ($4.66 \pm 12.21 \text{ W m}^{-2}$) in net radiation (Rn) is insignificant between the two ecosystems (Fig. S9 (a)) because the same meteorological forcings (mainly referring to downward shortwave and longwave radiation) were used for modelling. The averaged midday (i.e., 11:00 am) ground heat flux (G) accounted for 31 % (39 %) of Rn in the shrubs-grassland (grassland) ecosystem (Fig. S9). The observed importance of G was also reported in other semiarid and arid ecosystems with a dry soil surface and low vegetation coverage (Jia et al., 2016; Heusinkveld et al., 2004; Purdy et al., 2016; Kurc and Small, 2004).

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The Bowen ratio (i.e., H/LE) over two growing seasons was 71 % (67 %) of shrubs-grassland (grassland) ecosystems whereas, indicating that turbulent energy was dominated by LE in this semiarid region. The transition from LE to H dominance occurred from ~8:00 am to ~16:00 pm, during which the relatively high air temperature significantly affected H. Compared to the grassland ecosystem, the revegetated shrubs increased the H and LE. Similar findings have also been reported for a semiarid watershed in southeastern Arizona (Flerchinger et al., 1998), three semiarid ecosystems (cheatgrass, sagebrush and lodgepole pine) in the Snake River basin (Valayamkunnath et al., 2018) and a shrub-steppe ecosystem in Yanchi County (Gong et al.,

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2016). The consensus was that LE is positively correlated to the LAI, vegetation coverage and water availability. The H is positively correlated to the surface temperature.

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4.2.2 On gross primary productivity

Despite the underestimation of simulated GPP, the diurnal and monthly variations were in line with observations or simulations in other semiarid shrubs-grassland ecosystems in China (Jia et al., 2018; Du et al., 2021; Ma et al., 2020). It has commonly been concluded that revegetation enhanced carbon assimilation at the ecosystem level. Such an enhancement in this study (i.e., the difference in GPP between the two ecosystems) was more pronounced in 2019, which received more precipitation (Table S9). Besides, the significant carbon uptake in both ecosystems was noticed after each rainfall event (Fig. 9). This rainfall dependency of carbon flux is a representative characteristic in the semiarid regions, where the water and carbon cycles are tightly-coupled (Silva et al., 2017; Brümmer et al., 2008; Hastings et al., 2005; Eamus et al., 2013). Many studies reported that the shrublands are a stronger net carbon sink than the C3 grasslands (Eamus et al., 2013; Zhang et al., 2020a; Hastings et al., 2005; Petrie et al., 2015).

Interestingly, the GPP of the grassland ecosystem reached a plateau at ~10:00 am and then dropped at midday, particularly in July (Fig. S10 (a)). The likely cause is the midday depression phenomenon that high radiation and high surface temperature of leaf induce the saturation of photosynthesis, which followed by a decline of the stomatal conductance (Chen et al., 2014; Wang et al., 2019b; Deans et al., 2020). The simulations show the midday depression is less evident in the shrubs (Fig. S10), which is also evidenced by a field study on the transpiration characteristics of *Caragana intermedia* in the Mu Us Sandy Land (Zang et al., 2009).

4.2.3 On water fluxes

The simulated and observed SWC values at 10 cm soil depth agreed well during the calibration ($R^2=0.88$; $RMSE=0.01 \text{ m}^3 \text{ m}^{-3}$) and validation stage ($R^2=0.84$; $RMSE=0.01 \text{ m}^3 \text{ m}^{-3}$). Rather than the daily values of SWC, the instantaneous values at half-hour time steps were captured by the STEMMUS-SCOPE model, illustrating the timely SWC responses to rainfall events. This can facilitate the future investigation of the responses of plants (e.g., stomatal conductance and leaf water potential) to the water deficit (Liu and Shao, 2015; Fang et al., 2011). The rapid uptake of surface SWC that replenished by rainwater aligns with observations from *Caragana intermedia* plantations on the northeast Tibetan Plateau (Zhu and Wang, 2020).

Furthermore, our simulations indicated that SWC decreased within the 0-500 cm profile after revegetation. The most significant decrease in SWC occurred at soil depths of 35-280 cm, corresponding to the evident increase in RWU. Both phenomena were strongly associated with the root distribution and rooting depth of the shrubs. The shrubs extracted more soil

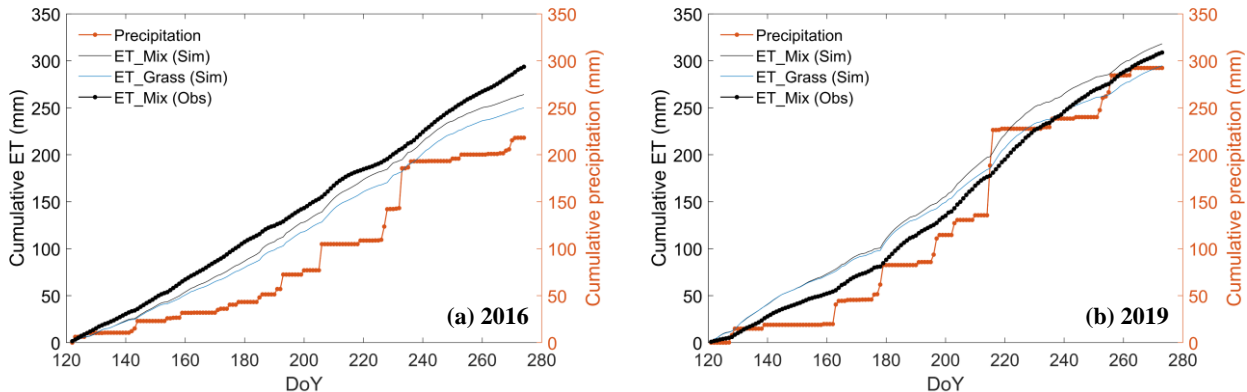
440 water than grasses, especially water from the deep soil. Moreover, such extraction was more intense in the drought year. Similarly, the simulations from the SHAW model suggested that SWC under the *Caragana Korshinskii* decreased within the 1.0-4.0 m profile and the SWC was depleted from deeper soil with the development of dry soil layers below 1.0 m (Liu and Shao, 2015). Our findings are also supported by field investigations on the root water uptake patterns of shrubs in the semiarid steppe of northern China, using stable isotopes technique or installing the sensors in multiple soil layers (Wang et al., 2018; 445 Zhu and Wang, 2020; Zhang et al., 2020b; Jia et al., 2012; Wang et al., 2019a; Jian et al., 2015). These experiments illustrated that the shrubs with a deep-rooted system, such as *Caragana* species, can flexibly switch their water source to deeper soil layers when the soil water in the shallow layers is depleted.

The seasonal evapotranspiration (ET) of the shrubs-grassland ecosystem was higher than that of the grassland ecosystem, 450 although the evaporation of the shrubs-grassland ecosystem was lower (Fig. 8 and Table S8). The revegetated shrubs have increased the LAI, directly diminishing the energy that reached the soil surface and thus decreasing evaporation. In this sense, revegetation increased the ecosystem ET mainly with the increasing transpiration (i.e., root water uptake). Our conclusions are qualitatively consistent with other studies in the same study area (Du et al., 2021; Dan et al., 2020), a shrub-encroached steppe ecosystem in Inner Mongolia (Wang et al., 2018), and a modeling study on the water-energy balance of shrubland-interspace 455 in Yanchi County (Gong et al., 2016).

An interesting finding is that the shrubs-grassland ecosystem presented a higher ratio between transpiration and evapotranspiration (T/ET) in the *dry* year 2016 (48 %) than that in *normal* year 2019 (41 %) (Table S8). This observation implies that the surplus rainwater in the wetter year was not absorbed effectively by roots but was evaporated or recharged 460 deep soils (> 300 cm) (Kurc and Small, 2007, 2004; Gao et al., 2023). Similarly, Chen et al. (2014) found that concentrated rainfall events did not induce a significant increase in transpiration of two revegetation species unless the rainwater could infiltrate the deep soils. At a semiarid loess site vegetated by apple trees in China's Loess Plateau, Gao et al. (2023) also observed a higher T/ET ratio during the extremely dry years as compared to extremely wet years; they also noted that the relationship between T/ET and annual precipitation could be complex (either independent or negatively correlated) and may 465 be influenced by soil characteristics. The intricate interplay between T/ET and annual precipitation of thick and dry loess soil profiles in the context of climate change presents a challenging topic.

4.3 Will the revegetated shrubs lead to soil water depletion?

For the grassland ecosystem, the ET partitioning, SWC at 10 cm and 100 cm depth and GPP remained relatively stable across two years. Planting shrubs, however, decreased the SWC and increased the water loss, leading us to hypothesize that the water consumption in shrubs-grassland ecosystem might reach its limit. To examine this assumption, we further compared the cumulative rainfall and ET in Fig. 11.



475 **Figure 11. Comparison of cumulative precipitation and cumulative Evapotranspiration (ET) of two ecosystems over the growing season in year (a) 2016 (dry) and (b) 2019 (normal), where the ‘ET_Mix (Sim)’ and ‘ET_Grass (Sim)’ are the simulated ET for shrubs-grassland and grassland ecosystem, respectively; and the ‘ET_Mix (Obs)’ is the observed ET (i.e., LE) from EC tower.**

The year 2016 witnessed an excess of 46 mm (76 mm) in cumulative simulated (observed) ET over the cumulative precipitation at the end of growing season (Fig. 11a). This observation points to an additional water source needed for ET, which can only be attributed to water transpired from deeper soil layers. In the wetter year 2019 (Fig. 11b), the rainwater replenishment seems enough to sustain the growth of grasses but still not enough for the shrubs-grassland ecosystem. Cumulative simulated (observed) ET of shrubs-grassland exceeded the precipitation by 26 mm (16 mm) in 2019. In comparison, the greater excess of ET over precipitation in 2016 provides evidence that drought conditions could result in a more significant depletion of water from deeper soil layers. Such excessive water consumption could lead to soil desiccation as reported in other semiarid regions with revegetation practices in China Loess Plateau (see supporting information Table S1 of Zhang et al., 2018). Nonetheless, our conclusions regarding water consumption exceeding ecosystem water supply are tentative, as they are based on simulations spanning only two growing seasons. Corroborating this, the whole year data in 2019 indicates that received rainfall (358.10 mm) was less than that of the ET from shrubland (400.15 mm) while was comparable to that from grassland (353.17 mm) (Dan, 2020). Long-term scenario simulations are necessary to ascertain the environmental threshold of exploitable water resources in the study area.

5 Conclusion

490 To understand the effects of revegetation on eco-hydrological processes of a desert steppe in northwestern China, we simulated the energy, water and carbon fluxes during May-September in 2016 (*dry*) and 2019 (*normal*), for a shrubs-grassland scenario and a grassland scenario, respectively. Simulations for two land covers were driven by respective LAI time-series and plant traits parameters in the STEMMUS-SCOPE model. Simulated fluxes based on half-hourly time steps agreed well with the measured trends. In particular, the model can simulate the soil water content accurately and capture its diurnal and daily
495 dynamics. According to the results of comparison between the two scenarios, the revegetation practices in the study area (i) increased the latent heat flux and sensible heat flux and decreased the ground heat flux, in which the latent heat flux dominated the energy partitioning; (ii) promoted gross primary productivity, which was highly responsive to rainfall availability; (iii) decreased soil water content at 0-500 cm soil depth (especially 35-280 cm) via root water uptake, which was more pronounced during the drier year; (iv) aggravated the water consumption of ecosystem with the decrease in soil water and remarkable
500 increase in root water uptake and transpiration. Moreover, revegetated shrubs appear to have a potential disruption in the water balance, manifested by greater evapotranspiration than received precipitation in two growing seasons. Future revegetation practices should consider the sustainable limits of ecosystems to avoid soil water depletion, which risks triggering the imbalance of the tightly-coupled energy, water and carbon cycles in the arid and semiarid regions.

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Code and data availability. The input data, source code and output data of STEMMUS-SCOPE are available on Zenodo at
510 <https://doi.org/10.5281/zenodo.10095916>.

Author contributions. ET, Z.Su and YZ designed and performed the study. ET ran the simulations, analysed the data and prepared the original draft. LD, HW and CQ offered the field data. YW, Z.Song, DY and CvdT provided technical help in simulations and data analysis. All authors reviewed and edited the manuscript.
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Competing interests. The authors declare that they have no conflict of interest.

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