# Riparian Ecohydrology: Regulation of Water Flux from the Ground to the Atmosphere in the Middle Rio Grande, New Mexico

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**Abstract**. During the previous decade, the southwestern United States has faced declining water resources and escalating forest fires due to long-term regional drought. Competing demands for water resources require a careful accounting of the basin water budget. Water lost to the atmosphere through riparian evapotranspiration (ET) is believed to rank in the top third of water budget depletions. To better manage depletions in a large river system, patterns of riparian ET must be better understood. This paper provides a general overview of the ecological, hydrological, and atmospheric issues surrounding riparian ET in the Middle Rio Grande of New Mexico. Long-term measurements of ET, water table depth, and micrometeorological conditions have been made at sites dominated by native cottonwood (Populus deltoides) forests and non-native saltcedar (Tamarix chinensis) thickets along the Middle Rio Grande (MRG). Over periods longer than one week, groundwater and leaf area index (LAI) dynamics relate well with ET rates. Evapotranspiration from P. deltoides forests was unaffected by annual drought conditions in much of the MRG where the water table is maintained above three meters below the surface. Evapotranspiration from a dense T. chinensis thicket did not decline with increasing groundwater depth; instead, ET increased by 50%, from 6 mm/day to 9 mm/day, as the water table receded at nearly 7 cm/day. Leaf area index of the T. chinensis thicket likewise increased during groundwater decline. Leaf area index can be manipulated as well following removal of non-native species. When T. chinensis and non-native Russian olive (Elaeagnus angustifolia) were removed from a P. deltoides understory, water salvaged through reduced ET was 26 cm/yr in relation to ET measured at reference sites. To investigate correlates to short-term variations in ET, stepwise multiple linear regression was used to evaluate atmospheric conditions under which ET is elevated or depressed. At the P. deltoides-dominated sites, ET anomalies were positively correlated to net radiation (R<sub>n</sub>) and negatively correlated to sensible heat flux (H), cross-corridor wind speed (v), and along-corridor wind speed (u) ( $r^2 = 0.54$ ). At the *T. chinensis*dominated sites, ET anomalies were positively correlated with R<sub>n</sub>, u, the friction coefficient (u<sub>\*</sub>), and vapor pressure deficit (VPD) and were negatively correlated to surface humidity scale (q\*), daily high and low temperature, H, and precipitation ( $r^2 = 0.66$ ). Both *Tamarix* and *Populus* can transpire prodigious quantities of water when the conditions are as favorable. In the Middle Rio Grande, T. chinensis is preferentially found where summer flooding and cold air drainage occurs, and P. deltoides is preferentially located in locations with shallow groundwater within two meters of the surface.

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#### Introduction

Riparian habitats worldwide are emerging as intense political, economic, social, and ecological battlegrounds over limited water resources (Zube and Simcox, 1987; Jackson *et al.*, 2001). In arid and semiarid regions, freshwater resources are concentrated along riparian corridors. These oases support high biodiversity, enhanced ecosystem productivity, and agricultural, municipal, and industrial demands for water (Malanson, 1995). The focus of human activities along rivers changes stream hydrology and ecology: perennial streams become ephemeral, diversions lower the water table, and plant stress leads to loss of native vegetation and shifts in species composition from native to non-native species (Medina, 1990; Smith *et al.*, 1991).

Increasing density of non-native species, along with declining ecological importance of native species, has been described in nearly all river systems throughout the western United States (Howe and Chancellor, 1983; Snyder and Miller, 1993; Di Tomaso, 1998; Pataki *et al.*, 2005; Shafroth *et al.*, 2005). The two most common non-native plants in western U.S. river corridors are *Tamarix* L. (saltcedar), and *Elaeagnus angustifolia* (Russian olive) (Friedman *et al.*, 2005). *Tamarix* spp. are primarily *T. ramosissima* and closely related *T. chinensis* and hereafter referred to as *Tamarix*. Both *Tamarix* and *E. angustifolia* originated in Eurasia, and they represent the third and fourth most common species found in western U.S. riparian areas, after *Salix exigua* (sandbar or coyote willow) and *Populus deltoides* (plains cottonwood) (Friedman *et al.*, 2005). The range of *E. angustifolia* is still expanding, while *Tamarix* dominance appears to be reaching an equilibrium in which expansion of *Tamarix* in some locations is balanced by loss of *Tamarix* in other locations (Friedman *et al.*, 2005). Examples of native species maintaining or

recovering community dominance have been observed in the Rio Grande and Colorado River systems (Snyder and Miller, 1993; Cleverly *et al.*, 1997; Nagler *et al.*, 2005b).

Riparian ecosystems that are prone to invasion by non-native species are characterized by greater disturbance frequency and intensity, deeper water tables, greater fluctuation of water table depth, greater surface salinity, and flood timing outside of the regeneration window for native species (Smith *et al.*, 1991; Busch and Smith, 1993; Busch and Smith, 1995; Planty-Tabacchi *et al.*, 1996; Lite and Stromberg, 2005; Tiegs *et al.*, 2005). Characteristics of *Tamarrix* that contribute to its ecological success include fruit production extending across the growing season; greater reproductive output; tolerance of heat, salinity, drought, and flooding stresses; regeneration following fire; induced sediment accretion with resultant deepening of the water table; surface soil salinization; and profligate transpiration from both groundwater and soil water sources (Warren and Turner, 1975; Busch *et al.*, 1992; Sala *et al.*, 1996; Di Tomaso, 1998; Cleverly *et al.*, 2002). Extensive reviews of *Tamarix* and *E. angustifolia* invasion ecology have been provided by Di Tomaso (1998) and Katz and Shafroth (2003), respectively.

The potential for non-native vegetation to remove prolific quantities of water from the ground is of serious concern to water resource managers, although a great deal of controversy continues to exist regarding water use by *Tamarix* (Hughes, 1993; Shafroth *et al.*, 2005). This controversy is driven by estimates of *Tamarix* water use that vary wildly, depending upon habitat and upon how water use is measured or estimated (Hughes, 1993; Cleverly *et al.*, 2002; Shafroth *et al.*, 2005). This paper will present a case study in riparian evapotranspiration along the Middle Rio Grande (MRG), New Mexico, U.S.A. There were two main objectives of this study: (1) to determine the relative evapotranspirational water losses from non-native and native riparian species under a range of natural environmental variations (Cleverly *et al.*, 2002; Dahm

et al., 2002); and (2) to investigate short- and long-term patterns of variability in measured ET for evaluating potential water salvage of riparian restoration and vegetation management (Molles et al., 1998; Shafroth et al., 2005; Newman et al., In Review).

### **Evapotranspiration**

Evapotranspiration (ET) is the process by which water is transported from the surface to the atmosphere as water vapor. Evapotranspiration is the combined evaporation from soil, open water, and ice surfaces, along with transpiration from vegetation. In riparian areas, the presence of shallow groundwater provides an ample water supply, permitting greater annual ET rates than precipitation (PPT) (Dahm *et al.*, 2002; Huxman *et al.*, 2005). Because ET from most ecosystems within a large basin is limited by PPT, riparian corridors exist as a hot spot of water loss to the atmosphere within large arid- and semiarid basins (Devitt *et al.*, 1998; Kurc and Small, 2004; Huxman *et al.*, 2005). Soil evaporation is negligible under riparian canopies due to attenuation of radiation by vegetation (Saugier and Katerji, 1991; Wilson *et al.*, 2001). Riparian ecosystems are characterized by a high transpiration to evapotranspiration ratio in the southwestern United States (Kurc and Small, 2004; Huxman *et al.*, 2005; Newman *et al.*, In Review).

Water budgeting has long been used to integrate large-scale surface water and groundwater dynamics. In large river systems, uncertainty in estimating the numerous pathways through which water can enter or leave a basin requires the use of the best "guesses" for depletions such as riparian ET, groundwater-recharge, and surface water-groundwater interactions (Scott *et al.*, 2000; Cleverly *et al.*, 2002; Dahm *et al.*, 2002). Of these depletions, ET from phreatophytes such as *P. deltoides* and *S. exigua* represents most of the 20 to 50 percent of total long-term

water budget depletions that are ascribed to natural vegetation (Scott *et al.*, 2000; Dahm *et al.*, 2002).

Many methods used for estimating ET are based upon the application of the energy balance, or conservation of incoming and outgoing energy. Under steady state conditions, and assuming a closed system, energy entering the canopy will be balanced by energy leaving the canopy:

$$R_{n} = \lambda E + H + G, \qquad (1)$$

where  $R_n$  is the net radiation,  $\lambda E$  is latent heat flux, H is sensible heat flux, and G is the ground heat flux (W m<sup>-2</sup>). Net radiation is the difference between downward- and upward-directed radiative energy flux:

$$R_n = (Q_{L\downarrow} + Q_{S\downarrow}) - (Q_{L\uparrow} + Q_{S\uparrow}), \tag{2}$$

where Q is radiative energy flux, L and S represent long-wave (i.e., thermal) and shortwave (i.e., solar) radiation, respectively. The direction of the arrow indicates whether radiation is directed down into the canopy or up away from the canopy. By convention,  $R_n$  is positive when directed toward the surface, and the remaining terms are positive when directed away from the surface. Latent heat flux represents the energy absorbed or released when water changes phase. Ground heat flux is the heat conducted through the soil surface, and H is the convective energy flux generated by atmospheric temperature gradients.

The Bowen ratio energy balance (BREB) method, for example, has been used for decades to estimate *Tamarix* ET (Gay and Fritschen, 1979; Devitt *et al.*, 1998). This method utilizes measurements of  $R_n$ , G, and the Bowen ratio,  $\beta$ , to determine  $\lambda E$ :

$$\beta = \frac{H}{\lambda E} = \frac{C_P}{\lambda \varepsilon} \frac{(T_2 - T_1)}{(\chi_2 - \chi_1)},\tag{3}$$

where  $C_p$  is the heat capacity of air,  $\lambda$  is the latent heat of vaporization, T is temperature,  $\epsilon$  is the ratio of the molecular weights of water and dry air, and  $\chi$  is the mole fraction of the partial pressure of water vapor. Temperature and humidity are each measured at heights two heights above the canopy where the subscript 2 refers to the upper measurement. Latent heat flux is determined as the residual of the energy balance (Eq. 1):

$$\lambda E = \frac{R_n - G}{1 + \beta} \,. \tag{4}$$

Bowen ratio energy balance methods have been applied successfully in wetlands with adequate fetch and a homogeneous canopy (Drexler *et al.*, 2004). Even when these conditions are met, however, energy balance estimates of ET can be confounded by sensible heat advection, which is the horizontal movement of momentum, energy, or other scalar quantities across a heterogeneous landscape (Itier *et al.*, 1994; McAneney *et al.*, 1994; Devitt *et al.*, 1998; Cooper *et al.*, 2000; Drexler *et al.*, 2004). Advection occurs when wet surfaces, such as wetlands or irrigated ecosystems, are located adjacent to dry surfaces such as arid and semi-arid rangeland ecosystems. Sensible heat advection into riparian forests is most easily identified when the evaporative fraction, which is the ratio  $\lambda E:R_n$ , is greater than unity (Devitt *et al.* 1998).

Three-dimensional sonic eddy covariance (3SEC) is the benchmark method for measuring fluxes over tall vegetation, under advection, and in complex terrain is (Brunet *et al.*, 1994; McAneney *et al.*, 1994; Simpson *et al.*, 1998; Drexler *et al.*, 2004). One advantage of the 3SEC system is that ET is measured directly, rather than estimated as the residual of the energy balance (Eq. 4). Thus, 3SEC is the only method that provides the means of estimating energy balance closure error as a self-check for accuracy (Twine *et al.*, 2000; Drexler *et al.*, 2004).

The core instruments in 3SEC systems are a 3-D sonic anemometer to measure wind speed in three dimensions, a krypton hygrometer or an infrared gas analyzer (IRGA) to measure humidity, a thermocouple with a very fine-wire junction (0.0127 mm diameter type E chromel–constantan) to avoid signal retention in high frequency measurements, and an IRGA to measure CO<sub>2</sub> concentration. Each of these instruments make measurements at a frequency determined through spectral decomposition of sample time series taken at 20 Hz. A sampling frequency is chosen in the high-frequency spectral gap, usually between 5 and 20 Hz (Stull, 1988).

Latent heat flux  $(\lambda E)$ , or the heat absorbed by evaporation, is computed from these measurements as a function of the covariance between vertical wind speed and humidity:

$$\lambda E = \lambda \overline{w'q'} = \lambda \sum_{i=1}^{n} \frac{(w_i - \overline{w})(q_i - \overline{q})}{n} = \lambda \text{Cov}(wq) = \lambda \rho_w \text{ET},$$
 (5)

where  $\lambda$  is the latent heat of vaporization, w' is the instantaneous deviation from mean vertical wind speed (i.e., w' = w<sub>i</sub> -  $\overline{w}$ ), and q' is the instantaneous deviation from average water vapor density. The time period for averaging, n, is chosen from balancing the spectral gap with minimizing trending in very low frequencies. Covariance periods are typically 15 to 40 minutes long. Sensible heat flux (H, heat transfer due to vertical atmospheric temperature gradients) is computed likewise as a function of the covariance between vertical wind speed and temperature:

$$H = C_p \rho_a \overline{w'T'} = C_p \rho_a \sum_{i=1}^n \frac{(w_i - \overline{w})(T_i - \overline{T})}{n} = C_p \rho_a \text{Cov}(wT),$$
 (6)

where  $\rho_a$  is density of wet air, and T' is the instantaneous deviation of temperature from mean air temperature.

Before ET is computed from LE, various standard corrections are applied to incorporate the physical realities within which these measurements are made. These corrections include coordinate rotation to align the wind vector with the sonic anemometer (Wesely, 1970),

corrections developed from frequency response relationships which incorporate sensor line averaging and separation (Massman, 2000; Massman, 2001), correction to account for flux effects on vapor density measurements as opposed to mixing ratio measurements (Webb *et al.*, 1980), and krypton hygrometer oxygen and second-order humidity effects on vapor density measurements. Lastly, closure of the energy balance is forced into thermodynamic equilibrium by adding closure energy into H and  $\lambda E$  in relation to the measured Bowen ratio (Twine *et al.*, 2000; Cleverly *et al.*, 2002).

Measurement of turbulent fluxes via 3SEC involves measuring flux variables from a tower or an aircraft. When measurements made from a tower, the minimum spacing between a canopy and the sensors is dictated by the necessity to make measurements above the roughness sublayer. The top of the roughness sublayer is equal to the distance of the aerodynamic roughness length,  $z_0$ , above the canopy,  $h_c$  (Nakamura and Mahrt, 2001). The upper limit for sensor placement in riparian corridors is chosen to remain below the top of the surface layer and to minimize the measurement footprint (Cooper *et al.*, 2000; Cooper *et al.*, 2003b). The top of the surface layer is defined in terms of Monin-Obukhov theory, which describes the physics of the atmospheric surface layer (Stull, 1988). Atmospheric stability imposed by advection is associated with vertical thermal stratification and compression of the surface layer to the lower few meters above the canopy (Cooper *et al.*, 2003b).

The planetary boundary layer is that lower portion of the atmosphere where turbulence dominates vertical transport and friction creates a strong drag on near-surface winds. The surface layer makes up the bottom 10% of the boundary layer (Stull, 1988), were surface drag creates a near-logarithmic wind speed profile. The vertical gradient in horizontal wind speed causes deformation of the air column and generates turbulence that can be characterized by

Reynolds stress,  $\tau_{Reynolds}$ , which is equal to the total vertical flux of horizontal momentum in the surface layer (Stull, 1988):

$$\left|\tau_{\text{Reynolds}}\right| = \overline{\rho} \left[ \left( u'w' \right)^2 + \left( v'w' \right)^2 \right]^{1/2} \tag{7}$$

where  $\rho$  is density of air, and u, v, and w are orthogonal wind speeds in two horizontal and one vertical dimensions, respectively. The velocity scale across which Reynolds stresses are propagated is the friction coefficient, u\* (Stull, 1988):

$$u_* = \frac{\left|\tau_{\text{Reynolds}}\right|}{\rho} = \left[\left(u'w'\right)^2 + \left(v'w'\right)^2\right]^{1/2} \tag{8}$$

The friction coefficient is important in describing the magnitude of turbulence in the surface layer, and that prominence is central to the body of the relationships known as Monin-Obukhov (M-O) similarity theory. M-O similarity theory, also known as surface layer similarity theory, is a body of empirical relationships that characterize turbulence in the surface layer. Other M-O scales useful for characterizing the surface layer include the humidity scale,  $q_*^{SL}$ , the temperature scale,  $\theta_*^{SL}$ , and the Obukhov length, L (Stull, 1988):

$$q_* = \frac{-\overline{w'q'}}{u_*},\tag{9}$$

$$\theta_* = \frac{-\overline{w'T'}}{u_*} \,. \tag{10}$$

$$L = \frac{-\overline{T}u_*^2}{kg\theta_*} \tag{11}$$

where k is von Kármán's constant and g is the acceleration due gravity. When L is zero, the surface layer is neutrally stratified. When negative, the surface layer is unstably stratified, and the temperature profile in the surface layer is stable when L is positive.

The measurement footprint represents the upwind distance that is incorporated into measurements made by instruments on the tower. The footprint increases with height above the canopy and with wind speed (Rannik *et al.*, 2000). In riparian forests along the Middle Rio Grande and the San Pedro River, the footprint for sensors mounted 2.5-m above *Tamarix* may be as little as 40 to 60 m under ideal conditions, or up to 200 m over *Populus fremontii* (Fremont cottonwood) and *Sporobolus wrightii* (sacaton grass) (Cooper *et al.*, 2000; Cooper *et al.*, 2003b).

#### The Middle Rio Grande

The Middle Rio Grande is located in New Mexico, U.S.A. and is defined to coincide with the New Mexico delivery obligation supply index gauges near Otowi NM and at Elephant Butte Dam, set forth in the Rio Grande compact (revised 1948). The MRG has three tributaries within the basin: the Jemez River, the Rio Puerco, and the Rio Salado (Fig. 1). The Rio Puerco and Rio Salado contribute flood flow to the MRG during summer monsoon periods. Three dams divert water from the Rio Grande to support irrigated agriculture (Fig. 1): the angostura diversion dam downstream of the confluence between the Rio Grande and the Jemez River, the Isleta pueblo diversion dam downstream of Albuquerque NM, and the San Acacia diversion dam downstream of the confluence of the Rio Grande with the Rio Salado. Three more dams create reservoirs in the MRG basin (Fig. 1): at Cochiti Pueblo, at Jemez Pueblo, and at Elephant Butte near Truth or Consequences NM. Elephant Butte Lake is the only reservoir with significant storage capacity, and the remaining reservoirs are principally operated for flood control. Mountainous terrain, identified in a 100-m resolution digital elevation model, was identified in the northern and southern portions of the MRG, while the Albuquerque and Belen reaches are overlooked by mesas (Fig. 1).

Flood control was promoted by the addition of riverside drains and levees during the early to mid 20<sup>th</sup> century. Exceptionally high flows in the river are contained within undeveloped riparian zones through the construction of levees. Adjacent to this levee system are drains that were excavated to an elevation below river channel bottom. These drains were placed to generate a hydrologic gradient away from the valley bottom, thus curtailing flooding beyond the levees. Excess water applied in irrigation is leached into the drain system.

One further significant hydrologic feature is the City of Albuquerque's wastewater reclamation plant. This plant was originally established in 1962 and contributes ca. 40% of the tributary inflow to the MRG, not including trans-basin transfers and tributary inflow in the upper Rio Grande basin above Otowi (Cleverly *et al.*, 2002; Dahm *et al.*, 2002). The origin of this water is pumped from the deep paleoaquifer at stations throughout the Albuquerque sub basin, providing long-term water resources to the MRG.

In arid and semi-arid regions, water loss to the atmosphere in riparian corridors dominate basin water budgets, representing well over 90% of Middle Rio Grande (MRG) depletions due to open water evaporation, soil evaporation, transpiration, and irrigated agriculture (Cleverly *et al.*, 2002; Dahm *et al.*, 2002). Evapotranspiration from riparian vegetation has been estimated to represent 30% of the total depletions in the MRG water budget.

Since the early twentieth century, riparian vegetation density and cover has been slowly increasing in all reaches of the Upper and Middle Rio Grande due to anthropogenic modifications of the MRG's hydrology (Everitt, 1998). On the Rio Grande upstream of its confluence with the Rio Puerco, *P. deltoides* cover has been increasing (Snyder and Miller, 1993). By the 1960's, woody vegetation cover reached a maximum in the MRG at Albuquerque, where *P. deltoides* represented nearly all of the 75% ground cover (Campbell and Dick-Peddie,

1964). Downstream of the Rio Grande's confluence with the Rio Puerco, *Tamarix* thickets and woodlands have expanded the area covered by woody vegetation while *P. deltoides* cover has decreased (Howe and Knopf, 1991; Everitt, 1998) (Fig. 1).

Currently, dense monocultures of non-native *Tamarix chinensis* Lour. (saltcedar) dominate the floodplain downstream of the Rio Grande's confluence with the Rio Puerco (Fig. 1). These thickets cover 100% of the land surface between riverside levees. In contrast, reaches of the Rio Grande upstream of the Rio Puerco's confluence are fully covered by a dense *Populus deltoides* ssp. *wislizeni* (S. Watson) Eckenw. (Rio Grande cottonwood) forests. In many locations, these *P. deltoides* forests include a dense understory of non-native species, predominately *Elaeagnus angustifolia* L. (Russian olive) and *Tamarix* (Fig. 1).

## Study area and methods

Four sites were chosen in 1998. Two locations of large extent were identified in *P. deltoides* forests, one with a short (i.e., less than three years) interflood interval, and the other with a long (i.e., greater than 40 years) interflood interval. Two *Tamarix* stands were likewise chosen with short and long interflood intervals (Molles *et al.*, 1998). The northernmost site, located in the south valley of Albuquerque, New Mexico, hosted a sparse *P. deltoides* forest (122 trees ha<sup>-1</sup>) and a very dense understory thicket of *Tamarix* and *E. angustifolia* until 2004, when the understory was removed as part of a citywide restoration project (Fig. 1). The other *P. deltoides*-dominated site, located near the towns of Belen and Rio Communities NM, regularly receives flooding and supports a high-density forest (278 trees ha<sup>-1</sup>) with predominantly native understory species. In November 1999, a 25-m tower was established to measure ET in each of these *P. deltoides* forests.

The northernmost *Tamarix*-dominated site is located at the Sevilleta National Wildlife Refuge (NWR), home of the Sevilleta Long Term Ecological Research (LTER) program. This site is located between the confluence of the Rio Salado and the San Acacia diversion structure, and the vegetation is a mosaic of *Tamarix* thickets, *D. spicata* meadows, and various halophytes like *Prosopis pubescens* Benth. (screwbean mesquite) and *Atriplex* L. spp (saltbush). A 10-m eddy covariance tower was established at this site in March 1999. The second *Tamarix*-dominated site is located at Bosque del Apache NWR. This site is frequently flooded and contains a dense (15,000 shrubs ha<sup>-1</sup>), monospecific *Tamarix* thicket. The U.S. Bureau of Reclamation and New Mexico State University established a 15-m tower in this thicket in 1998.

Three-dimensional eddy covariance system were mounted on the towers, 2 to 2.5 meters above the canopy (Cleverly *et al.*, 2002; Dahm *et al.*, 2002). The measurement period was 10 Hz and the covariance period was 30 minutes. Additional energy flux measurements were made at 1 Hz and averaged over 30 minutes. Net radiation was measured using annually cross-calibrated REBS Q7.1 net radiometers. Ground heat flux was measured using Campbell Scientific, Inc. (CSI) ground heat flux plates, corrected for heat storage in the 8 cm of soil above the plate using soil temperature (CSI TCAV) and soil water content (CSI CS616) measurements. Average energy balance closure for all systems in all years, determined on a daily basis, was 78%, and the Bowen ratio was near zero during the growing season (Cleverly *et al.*, 2002). These are values typical of eddy covariance and riparian corridors, respectively (Devitt *et al.*, 1998; Wilson *et al.*, 2002).

Complementary measurements made from each tower include standard weather station conditions (i.e., precipitation, temperature, relative humidity, horizontal wind speed and direction; Campbell Scientific, Inc., Logan, UT). Measurements of vapor pressure were used to

calculate vapor pressure deficit (VPD) as the difference between atmospheric vapor pressure (e<sub>a</sub>, kPa) and saturation vapor pressure (e<sub>s</sub>, kPa). This difference represents the vapor pressure gradient from the saturated tissues within a leaf to the free atmosphere.

A network of five shallow groundwater wells was placed at each site, with one well near the tower and four additional wells in the four cardinal directions, 40-m from the central well. Three of the wells at each of the sites were equipped with automated pressure transducers, measuring water level every 30 minutes (EEI, Las Cruces, NM). In 2003, all five wells at each site were instrumented with pressure transducers, with a subset of those also measuring groundwater temperature (Solinst Canada Ltd, Georgetown, Ontario).

Plant area index (PAI), which is the projected area of leaves and stems per unit ground area, was measured from below relatively undisturbed vegetation adjacent to each of five groundwater wells using the LAI2000 (Li-cor, Inc., Lincoln, NE). Two measurements above the canopy and five measurements below the canopy were made in each plot. The first and second plots were double-sampled to track changes in measured PAI with changing light conditions. Measurements were made monthly during the growing season, as well as once during the winter of 2001-2002 to estimate stem area index (SAI). Retention by *P. deltoides* of senescent leaves through the winter confounds the winter measurement from *P. deltoides* forests. Measurements were made either during 45 minutes between sundown and dark or, weather permitting, under cloud cover. Measurements of clear sky conditions were made from a ladder or from the tower at *Tamarix* sites and from the nearby levee at *P. deltoides* sites because the vegetation is taller than 20-m.

Plant area index is equal to the sum of the projected LAI and SAI, less the area of overlap between stems and leaves. In broadleaf species such as *P. deltoides*, overlap between stem area

and leaf area is the same as SAI, thus PAI equals projected LAI. (Gower and Norman, 1991; Chen and Black, 1992; Chen and Cihlar, 1995; Chen and Cihlar, 1996). Incorporation of leaf geometry was considered in three leaf types: broadleaf tree (i.e., *P. deltoides* and *E. angustifolia*), flat leaf shrubs (i.e., *Salix exigua*), and cylindrically-leaved shrubs (i.e., *Tamarix*). Leaf area index of the three-dimensional leaf surface was determined for each leaf type as:

$$LAI(broadleaf)_{total} = 2 \bullet PAI$$
 (12)

$$LAI(flat)_{total} = 2 \bullet (PAI - SAI)$$
(13)

$$LAI(cylindrical)_{total} = \pi \bullet (PAI - SAI)$$
(14)

Stepwise multiple linear regression (proc GLM, Statistical Analysis Software, SAS Institute, Inc., Cary, NC) was used to evaluate effect of micrometeorological conditions on daily ET rates (Nagler *et al.*, 2005a). A suite of micrometeorological variables was chosen for the analysis representing surface layer aerodynamics, energy fluxes, and standard weather phenomena (Table 2). Because temporal autocorrelation in micrometeorological time series data violate the critical regression assumption of independent and identically distributed (i.i.d.) measurements, anomaly analysis was chosen to resample the time series and thereby eliminate i.i.d. restrictions (Higgins *et al.*, 1997). Evapotranspiration anomalies are identified as having a value greater or lesser than one standard deviation from average, computed as the 9-day running average,  $\overline{ET_{9-day}}$ . These n=43±3 anomalies per year from each tower are then used as the dependent variable in both forward and reverse steps. This analysis indicates which micrometeorological conditions are related to driving daily ET above or below the long-term average.

## **Drought and Groundwater (slow dynamics)**

In the year 2000, the *P. deltoides* forest with a non-native understory had the highest annual ET (Albuquerque) (Table 1). The lowest ET in every year was generated by the xeroriparian *Tamarix* woodland at the Sevilleta NWR (Table 1). This xeroriparian *Tamarix* woodland was also distinguished by having the highest ET:LAI ratio of all the sites (Table 1). The vegetation in this location is predominantly a mosaic of *Tamarix* and *D. spicata*. *Prosopis pubescens* (screwbean mesquite) and *Atriplex* spp. are also present at the Sevilleta site. Of the sites with intermediate annual ET, monospecific *Tamarix* (Bosque del Apache NWR) lost more water through ET than the *P. deltoides* forest containing a mostly native understory (Belen—Rio Communities) (Table 1). Average annual precipitation is 20 to 31 cm in the MRG (Dahm *et al.*, 2002), and the ET:PPT ratio was greater than unity at all sites in all years.

Removal of non-native species from the understory of a *P. deltoides* forest had a modest effect on forest ET. Evapotranspiration decreased by 9% following removal of non-native species between 2003 and 2004 (Table 1). However, ET at both of the *Tamarix*-dominated sites increased in 2004 by 12% over 2003 ET (Table 1). Assuming that the *Tamarix* sites are representing the reference conditions for ET water loss in the basin, the net decrease in ET due to removal of non-native species is 21%, a savings of 26 cm in the first year following clearing. While this value is consistent with earlier predictions, water salvage may be short-lived because *Tamarix* and *E. angustifolia* were allowed to regenerate during the 2005 growing season (data not shown) (Shafroth *et al.*, 2005). Greater reduction in ET is expected to follow from converting a monospecific saltcedar stand (e.g., Bosque del Apache NWR) into a saltgrass or meadow (Shafroth *et al.*, 2005).

Recurrent, extended drought in the southwestern United States, also referred to as "megadrought," is characterized decreased regional precipitation in both the winter and summer modes. Recently, research has shown long-term sea surface temperature anomalies to be related to these megadroughts, in which the Pacific decadal oscillation is in a cool phase (PDO-) and the Atlantic multi-decadal is in a warm phase (AMO+) (Gray *et al.*, 2003). Resultant decline of regional precipitation and snowpack is initially ameliorated in the riparian corridor by release from reservoir storage, maintaining surface flow and groundwater resources. Along the MRG, droughts have recurred every 20 to 70 years (Gray *et al.*, 2003).

Drought is expected to have an important recurring effect upon the MRG water budget, especially in naturally occurring processes like ET from riparian vegetation. Reduction in ET from *P. deltoides* stands was minimal, even during extreme drought (Table 1). The vegetation did not experience the full intensity of what has been considered a major drought because municipal wastewater maintained the water table within two meters of the surface at the Albuquerque site (Table 1 and Fig. 2). Likewise, groundwater dynamics follow a hydrograph of irrigation operations rather than a natural flow regime at the Belen site (data not shown). Flow in the riverside drain maintained the water table above 175 cm below the surface (Table 1). As long as the irrigation and wastewater treatment systems are in operation, ET depletions from *P. deltoides* forests remain undimished (Table 1).

Evapotranspiration and LAI declined in the *Tamarix* sites during drought (Table 1). Evapotranspiration at the Sevilleta NWR's *Tamarix* and *D. spicata* woodland tracked drought severity index values closely (Table 1). Bosque del Apache NWR's monospecific *Tamarix* thicket initially followed this trend, but ET and LAI began returning to pre–drought levels one year early, in 2003 (Table 1). Groundwater levels also declined with drought at the *Tamarix* 

sites, especially during the late summer (Table 1 and Fig. 2). Groundwater levels and annual variability had not begun to return to pre-drought patterns in 2004, the first year following relief from meteorological drought (Table 1 and Fig. 2).

Evapotranspiration is often modeled as a function of groundwater depth (McDonald and Harbaugh, 1988). In the earliest conceptualization of this relationship, an elevation was defined above which ET is not limited by groundwater availability (i.e., ET<sub>surface</sub>). At some deeper elevation, ET drops to zero because the water table is too deep for roots to access (i.e., ET<sub>extinction</sub>). Surface and extinction elevations are species specific. In *P. deltoides*, for example, these depths are ET<sub>surface</sub> = 3-m and ET<sub>extinction</sub> = 5-m (Horton *et al.*, 2001). In fact, *P. deltoides* is well known for intolerance to groundwater recession, and crown dieback has been observed when the water table falls below three meters (Scott *et al.*, 1999; Rood *et al.*, 2000; Cooper *et al.*, 2003a). *P. deltoides*-dominated forests are typically found in places with a shallow and stable water table (Table 1 and Fig. 2).

While ET<sub>surface</sub> and ET<sub>extinction</sub> are well defined in *P. deltoides*, depth to groundwater has not been found to restrict *Tamarix* ET, even when the water table is 10 m (Horton *et al.*, 2001) or 25 m (Gries *et al.*, 2003) below the surface. In the latter study, water storage in sand dunes sustained transpiration because *Tamarix* is a facultative phreatophyte capable of exploiting soil water in addition to groundwater (Busch *et al.*, 1992; Smith *et al.*, 1998). *Tamarix* tend to occupy sites with groundwater depth variability (Lite and Stromberg, 2005), and that is true at Bosque del Apache NWR where water table fluctuates more than 350 cm per year (Fig. 2).

Water table depth was deepest in *Tamarix*-dominated sites (Table 1). In 2003, the water table at Bosque del Apache NWR fell below 3.5 meters for the first time in this study (Fig. 2). Groundwater decline was extreme that summer, falling as rapidly as seven cm day<sup>-1</sup> (Fig. 3) from

175 cm below surface in May to 360 cm in September (Fig. 2). While crown dieback occurred in a nearby *P. deltoides* forest (data not shown), a resurgence in *Tamarix* ET accompanied groundwater recession (Table 1 and Fig. 3). ET from this *Tamarix* thicket increased substantially within a few days following the initiation of groundwater decline. Recently, Naumburg et al (2005) proposed a mechanism that explains this counterintuitive finding.

For species like *Tamarix* that are not susceptible to moderate soil drying (Pockman and Sperry, 2000), a receding water table can increase the volume of aerated soil near field capacity (Naumburg *et al.*, 2005). Draining of the soil is slower than the potential root growth rates in *Tamarix*, and the partly drained soil can be easily exploited (Naumburg *et al.*, 2005). The initial delay results from the time required for the capillary fringe of the original water table ( $GW_0 = 175$  cm) to drain sufficiently to allow respiration and root growth below  $GW_0$ . Once this hypothetical taproot growth is initiated, water uptake by new root growth at depths greater than  $GW_0$  is augmented by continued fine root functioning at  $GW_0$ . Fine roots at the previous capillary fringe continue to function because *Tamarix*, unlike *P. deltoides*, is relatively tolerant of low xylem water potential (Pockman and Sperry, 2000).

A few days before 1 July, ET decreased and groundwater recession slowed (Fig. 3). Stand ET declined again at the end of the year, preceding the year's maximum water table depth (GW<sub>max</sub>) (Figs. 2 & 3). Decreased groundwater decline rate and ET near the beginning of July was caused by multi-day monsoon precipitation, during which net radiation and vapor pressure deficit (VPD) were minimized (data not shown). When that system had passed, groundwater decline and enhanced ET returned (Fig. 3).

Dependence upon the shallow aquifer may be related to phreatophytic status (Smith *et al.*, 1998). Thicket forming *Tamarix* spp. in the western U.S. are facultative phreatophyte using

water of an isotopic ratio intermediate between groundwater and precipitation sources (Busch *et al.*, 1992). In *Populus* spp., use of precipitation water sources is conditional upon access to groundwater; in locations with a deep water table or ephemeral access to a shallow water table, *Populus* spp. utilize both deep and shallow water sources (Stromberg and Patten, 1996; Snyder and Williams, 2000). Of the dominant species along the MRG, *Elaeagnus angustifolia* (Russian olive) is extremely varied in the habitats in which it thrives, but little has been documented regarding distributions where the groundwater is more than a few meters below the surface (Katz and Shafroth, 2003).

Average high and low temperature varied only slightly during between 2000 and 2004 (Table 1). The southern sites experience hotter days and colder nights than the northern sites (Table 1). This unusual pattern of minimum temperatures decreasing downstream can be explained by considering the distinct topographic features surrounding each site (Papadopoulos and Helmis, 1999). Katabatic winds, better known as cold air drainage, tends to be greatest in gullies and in mountainous terrain (Papadopoulos and Helmis, 1999; Soler *et al.*, 2001). These mountainous hillsides have slopes that increase in angle from the valley floor to mountain peaks. Because sunrise does not illuminate the entire east-facing slope at once, an inversion layer is formed just below the sunrise line. Katabatic flow continues below this inversion layer and anabatic flow begins above (Papadopoulos and Helmis, 1999). Because of the mountainous terrain near the downstream reaches of the MRG, cold air drainage into the *Tamarix* thickets maintains colder nighttime temperature (Table 1).

### Flooding and Micro-meteorology (rapid dynamics)

Limited flooding of two sites, Belen—Rio Communities and Bosque del Apache NWR, occurred during spring 2001 due to a one-day release of 115 m<sup>3</sup> s<sup>-1</sup> from Cochiti Reservoir by the

US Army Corps of Engineers (Fig. 4). The travel time of this flood pulse was 48 hours between these two sites (Fig. 4). At Bosque del Apache, floodwater flowed into the floodplain over the bank 0.5 to 1 km upstream of the tower. Surface flow traveled parallel to the river, in contrast to flooding near Rio Communities, where stagnant open water arose from seepage. Surface inundation was short-lived at both sites, and the water table peaked for a single day (Fig. 4).

During onsite flooding, groundwater levels in the monitoring wells failed to reach the surface at either flooded site. At the northern cottonwood-dominated site, the water table reached the surface in some locations but not others, averaging 55 cm below the average land (Fig. 4). Local topographical relief is 50 cm between the highest and lowest-lying wells, but these wells represent only a fraction of the total topographic relief. Surface flooding was likewise observed in depressions but not ridges within the active floodplain.

Total ET from this *P. deltoides* forest did not increase during or following inundation (Fig. 4). Net radiation reaching the forest floor through the canopy is minimal, generally between 50 and 100 W m<sup>-2</sup>, providing little energy for forest floor evaporation (Wilson *et al.*, 2001). At the same time, transpiration is not curtailed completely due to root hypoxia, for only a fraction of the active roots are likely to be under the water table.

Flooding in the monospecific *Tamarix* thicket was associated with elevated total ET rates (Fig. 4). Unlike the loamy sand underlying the *P. deltoides* forest, the soil at this site is dominated by loamy clay. The Rio Puerco is well known for generating a load of very fine sediments into the MRG (Phippen and Wohl, 2003; Gellis *et al.*, 2004). Fine sediments defloculate when wetted, creating an infiltration barrier. Therefore, the top meter of soil did not reach field capacity even though surface water was observed around all groundwater wells (Fig. 4).

Evapotranspiration rates were closely coupled to VPD at the *Tamarix* sites (Table 3 and Fig. 4). *Tamarix*' cylindrical leaves form a minimal boundary layer, and stomata respond rapidly to atmospheric conditions and thereby closely link transpiration to micro-meteorological conditions (Table 3) (Jarvis and McNaughton, 1986; Hollinger *et al.*, 1994; Meinzer *et al.*, 1997; Nagler *et al.*, 2005a). At the *P. deltoides* sites, coupling between VPD and ET was observed for a part of the year, but asynchrony between VPD and ET fluctuations was observed during flooding (Fig. 4). On average, short-term fluctuations in *P. deltoides* ET were not closely coupled to fluctuations in VPD (Table 3).

Coupling between ET and atmospheric conditions was closer at the *Tamarix* sites than at the *P. deltoides* sites (Table 3 and Fig. 5). Evapotranspiration from the *Tamarix* canopy was especially sensitive to aerodynamic characteristics of the surface layer (Table 3). Variations in  $u_*$  are observed under advection, when temperature inversions within the lower atmosphere constrain the extent of the surface layer (Cooper *et al.*, 2003b).

Precipitation decreases ET and H at the *Tamarix* sites through shading and the resulting reduction in  $R_n$  (Table 3). Because vegetation along the MRG are not water limited due to the close proximity of groundwater (Table 1 and Fig. 2), precipitation does not directly lead to changes in ET (Table 3), in contrast to the flush of leaf gas exchange that follows precipitation in drier riparian habitats (Huxman *et al.*, 2004). The interaction of daily temperature extremes also contributed to the micro-meteorological control of *Tamarix* ET (Tables 1 & 3) (Nagler *et al.*, 2005a).

Energy balance and horizontal wind direction explained slightly more than 50% of the daily variability in *P. deltoides* ET (Table 3 and Fig. 5). Sensible heat advection, carried on the wind tangential to the riparian corridor, and the development of a thick local boundary layer,

reduces the coupling between stomatal conductance, vapor pressure deficit, and transpiration (Table 3) (Jarvis and McNaughton, 1986). Sensible heat advection and surface layer dynamics had a stronger effect on *P. deltoides* ET in Albuquerque's south valley than in Belen (data not shown).

Modeled ET anomalies fit well with measured anomalies, especially at the *Tamarix* sites (Fig. 5). Residuals from the regression analysis were evenly distributed around the model line (Fig. 5), suggesting that the anomaly dataset is essentially free from autocorrelation bias (Nagler *et al.*, 2005a). For the most part, energy balance and aerodynamics are related to short-term fluctuations in riparian ET (Table 3) (Pieri and Fuchs, 1990). Variables that were included in the analysis, but which did not show any relationship to ET fluctuations, included depth to groundwater, G, and barometric pressure (Tables 2 & 3). Ground heat flux is near zero and does not vary (data not shown). Depth to groundwater does vary, but at a slower scale that could not be detected in the short-term analysis.

#### Conclusions

Evapotranspiration rates were compared among native and non-native riparian vegetation along the Middle Rio Grande in New Mexico. Evapotranspiration from xeroriparian shrubs, including non-native *Tamarix* and native *D. spicata*, was lower than ET from any other vegetation class (Table 1). Highest ET rates were measured from a multi-layered canopy site in which non-native *Tamarix* and *E. angustifolia* formed a thick understory with a *P. deltoides* forest (Table 1). However, the monospecific *Tamarix* and native *P. deltoides* forest without a dense non-native understory also support ET rates greater than 100 cm yr<sup>-1</sup>, which is three to five times annual precipitation in the basin (Table 1) (Dahm *et al.*, 2002). Energy balance closure

error was 78%, well within the 10%-30% range for eddy covariance systems (Twine *et al.*, 2000).

Reduction in ET following removal of the non-native understory from a *P. deltoides* canopy resulted in a modest 21% reduction in ET relative to the reference sites (Table 1) (Shafroth *et al.*, 2005). However, ET at the post-removal was lower than it had been in any other year at that site, it was the first year this site did not support higher ET rates than any other site. Water salvage will be transitory, however, should the understory be allowed to regenerate its non-native understory. To be economically feasible, an emphasis upon long-term success in restoration projects needs to be pursued.

Evapotranspiration rates responds to groundwater depth, stomatal coupling to atmospheric humidity, leaf area index, atmospheric surface layer conditions, and energy balance. The recent severe drought had little effect on native *P. deltoides* forests because water tables were artificially maintained to within two meters of the surface, but *Tamarix* water use and LAI were diminished under drought conditions (Table 1). However, water table decline during drought resulted in 50% higher ET rates from a dense, monospecific *Tamarix* thicket (Fig. 3). Evapotranspiration from *Tamarix* is closely coupled to atmospheric conditions rather than water table depth (Table 3 and Fig. 4). Evapotranspiration from *P. deltoides* is responsive to variations in energy balance and sensible heat advection rather than atmospheric humidity or temperature (Table 3) (Horton *et al.*, 2001).

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Table 1. Summary of drought severity, annual evapotranspiration (ET, cm yr $^{-1}$ ), temperature extremes, and groundwater depth. Average annual Palmer drought severity index (PDSI) scores are the average monthly values from New Mexico regions 2 (North-central New Mexico) and 5 (Middle Rio Grande) (http://www.ncdc.noaa.gov/oa/climate/onlineprod/drought/xmgrg3.html). More negative PDSI values indicate increasingly severe drought. Average daily temperature extremes, both nightly low (Tmin) and the daytime high (Tmax), were calculated as an average of the same 92 days in each year: 14-23~May, 13-16~June, 21~June-2~July, 11-12~July, 10-15~August, and 1~September-28~October. Maximum daily average groundwater depth (GWmax) was measured from a well within five meters of each tower.

	Year:	2000	2001	2002	2003	2004
Site	PDSI:	-0.04	1.01	-3.39	-3.94	-0.09
	ET (cm/yr):	134	125	127	125	114
Albuquerque	$T_{min} / T_{max}$	12.3 / 26.5	12.0 / 27.1	12.1 / 26.5	12.9 / 27.4	12.1 / 26.0
Populus deltoides*	LAI <sub>Total</sub>	$6.6 \pm 0.2$	$7.0 \pm 0.2$	$6.4 \pm 0.2$	$4.0 \pm 0.2$	
L	Gw <sub>max</sub> (cm)	(171)	(175)	(178)	(190)	(186)
	ET (cm/yr):	103	107	95	126	
<i>Belen</i>	$T_{min} / T_{max}$	12.4 / 27.2	11.2 / 27.9	11.7 / 27.5	12.2 / 28.9	
Populus deltoides	LAI <sub>Total</sub>	$5.0 \pm 0.2$	$6.4 \pm 0.2$	$6.0 \pm 0.2$	$5.2 \pm 0.2$	
L	Gw <sub>max</sub> (cm)	(1 <u>18</u> )	(139)	(1 <u>55</u> )	(168)	(163)
	ET (cm/yr):	86	83		69	
Sevilleta NWR	$T_{min} / T_{max}$	10.1 / 28.6	8.8 / 29.5	9.2 / 28.5	9.8 / 29.6	8.6 / 27.8
Tamarix chinensis**	LAI <sub>Total</sub>	$3.5 \pm 0.1$	$1.6 \pm 0.2$	$0.9 \pm 0.1$	$1.6 \pm 0.2$	
L	Gw <sub>max</sub> (cm)	(213)	(218)	(227)	(245)	(240)
	ET (cm/yr):		115	87	101	114
Bosque del Apache NMR	$T_{min} / T_{max}$	9.7 / 28.6	7.8 / 29.2	9.2 / 29.4	9.3 / 30.4	9.0 / 28.1
Tamarix chinensis	LAI <sub>Total</sub>	$4.7 \pm 0.1$	$5.0 \pm 0.1$	$2.5 \pm 0.1$	$4.1 \pm 0.1$	
	Gw <sub>max</sub> (cm)	(267)	(288)	(347)	(385)	(376)

<sup>\*</sup> dense non-native understory removed winter 2003/2004

<sup>\*\*</sup> mosaic with *Distchlis spicata*, *Prosopis pubescens*, and *Atriplex* spp.

Table 2. Variables evaluated in stepwise multiple regression. During each iteration, variables with a probability of a type II error > 0.1 (p > 0.1) was removed from the analysis.

Variable	Units	Definition					
Energy Balance:							
$R_n$	$W m^{-2}$	Daytime Net Radiation					
Н	$W m^{-2}$	Sensible Heat Flux					
G	W m <sup>-2</sup>	Ground Heat Flux					
Aerodynamics:							
u	m s <sup>-1</sup>	Daytime Windspeed along riparian corridor					
V	m s <sup>-1</sup>	Daytime Windspeed across (i.e., tangential to) riparian corridor					
WS	m s <sup>-1</sup>	Daytime 2-D Horizontal windspeed					
nu	m s <sup>-1</sup>	Nighttime windspeed along riparian corridor					
nv	m s <sup>-1</sup>	Nighttime windspeed across riparian corridor					
nws	m s <sup>-1</sup>	Nighttime 2-D Horizontal windspeed					
u∗	m s <sup>-1</sup>	Friction coefficient or velocity					
q*	$g_{water}  g_{air}^{-1}$	Humidity scale in the surface layer					
L	m	Obukhov Length					
Meteoro	Meteorological Conditions:						
GDD	°C	Growing Degree Days					
PPT	mm day <sup>-1</sup>	Precipitation					
q	9 <sub>water</sub> 9 <sub>air</sub> -1	Specific Humidity					
T <sub>max</sub>	°C	Daily High Temperature					
T <sub>min</sub>	°C	Daily Low Temperature					
Т	°C	Daily Average Temperature					
VPD	kPa	Vapor pressure deficit					
Р	kPa	Atmospheric Pressure					
Other:							
GW	cm	Depth to groundwater					

Table 3. Results of stepwise multiple regression model between evapotranspiration anomalies and micrometeorological conditions. ET anomalies were identified when daily ET was more than one standard deviation above or below the 9-day running average. For ease of interpretation, factors related to ET anomalies are separated into energy, aerodynamic, and meteorological conditions. See text and Table 2 for descriptions of the modeled dependent variables. Models for predicting ET anomalies are  $[ET_{anomaly} = -0.008~H + 0.02~R_n - 0.1~v - 0.09~(v~X~u)]~(r^2 = 0.54)$  in cottonwood forests and  $[ET_{anomaly} = 0.005~R_n + 0.08~u + 1.2~u* - 4.2~q* + 11.8~(u* X~q*) - 0.5~VPD - 0.01~(T_{max}~X~T_{min}) - 0.003~(PPT~X~H)~+ 0.001~(R_n~X~PPT)]~(r^2 = 0.66)$  for saltcedar stands.

Factor	Coefficient ± se	F	р					
Albuquerque and Belen — Rio Communities,								
Populus deltoides								
Model:	0.54	110.8	< 0.0001					
Energy Balan								
	$-0.008 \pm 0.002$							
$R_n$	0.02 ± 0.0008	388.1	< 0.0001					
Aerodynamics:								
V	-0.1 ± 0.06	5.8						
v X u	-0.09 ± 0.02	16.2	< 0.0001					
Sevilleta a	nd Bosque del Ap	ache N	WRs,					
Tamarix chinensis								
Model:	0.66	77.7	< 0.0001					
Energy Balance:								
$R_n$	$0.005 \pm 0.0005$	83.7	< 0.0001					
Aerodynamics:								
u	$0.08 \pm 0.03$	7.5	0.007					
U∗	$1.2 \pm 0.3$	12.9	0.0004					
q∗	$-4.2 \pm 0.6$	50.2	< 0.0001					
u∗ X q∗	11.8 ± 4.3	7.4	0.007					
Meteorologica	al Conditions:							
VPD	$0.5 \pm 0.07$	43.0	< 0.0001					
$T_{max}XT_{min}$	-0.01 ± 0.003	9.8	0.002					
Interacting Effects:								
	$-0.003 \pm 0.0005$	24.3	< 0.0001					
R <sub>n</sub> X PPT	0.001 ± 0.0003	18.4	< 0.0001					

# **Figure Legends**

Figure 1. Map of the Middle Rio Grande in New Mexico, USA. Locations of local cities are marked in the inset map. The Middle Rio Grande, originating in Otowi NM and discharging from Elephant Butte Reservoir, is dominated in the north by *Populus deltoides* ssp. *wislizeni* (Rio Grande cottonwood) and by *Tamarix chinensis* (saltcedar) in the southern reaches. Locations of nearby mountain ranges are illustrated, along with diversion dams, reservoirs, and tributaries. Four sites with tower-based eddy covariance systems are listed along with site locations and local riparian vegetation.

Figure 2. Daily average depth to groundwater (cm) in Albuquerque's South Valley (ABQ) and at Bosque del Apache NWR (BDA). Pressure transducer measurements were logged once every 30-min from within five meters of the tower.

Figure 3. Temporal pattern of mean changes in groundwater depth and mean Tamarix chinensis evapotranspiration (ET) in 2003 at Bosque del Apache National Wildlife Refuge. Change in groundwater depth is the difference between mean water table depth between two consecutive days,  $dGW/dt = GW_{t-1} - GW_t$  where t is a discrete day. Negative values indicate a falling water table. Groundwater depth change and ET were smoothed with a 9-day running average. Refer to Figure 2 for daily depth to water table. The first day of each month is highlighted, and arrows follow the progression of the growing season.

Figure 4. Time series of daily evapotranspiration (ET), vapor pressure deficit (VPD), and depth to groundwater. Floodwaters were present at the Belen — Rio Communities *Populus deltoides* ssp. *wislizeni* (A) and Bosque del Apache *Tamarix chinensis* (B) sites on May 24 and May 26, 2001, respectively. The arrows in panel A and the first pair of arrows in panel B point

to the day of inundation. The two following pairs of arrows in panel B point to days of following ET maxima.

Figure 5. Measured and predicted evapotranspiration anomalies. Predicted anomalies were defined as the solution to the best-fit stepwise multiple linear regression model (Table 3).

Figure 1:

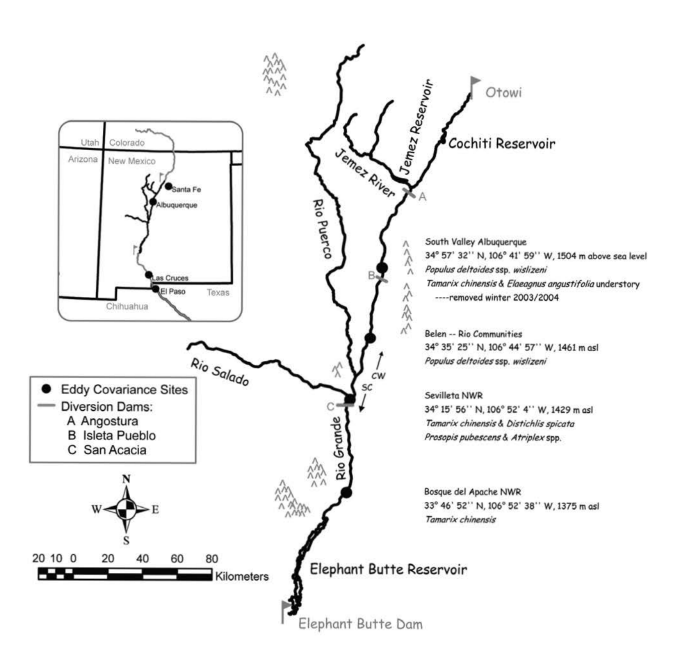


Figure 2:

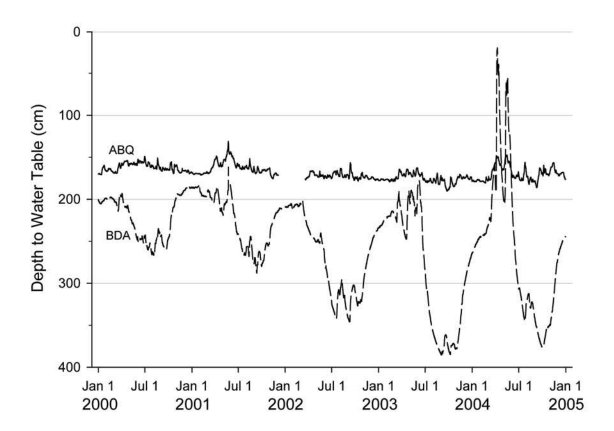


Figure 3:

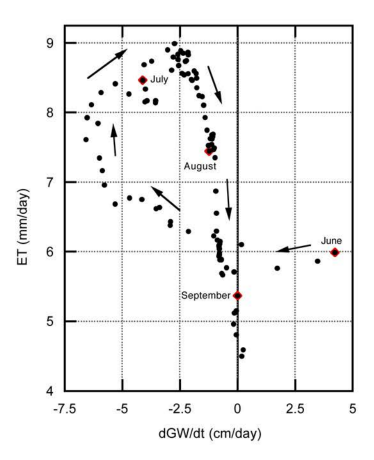


Figure 4:

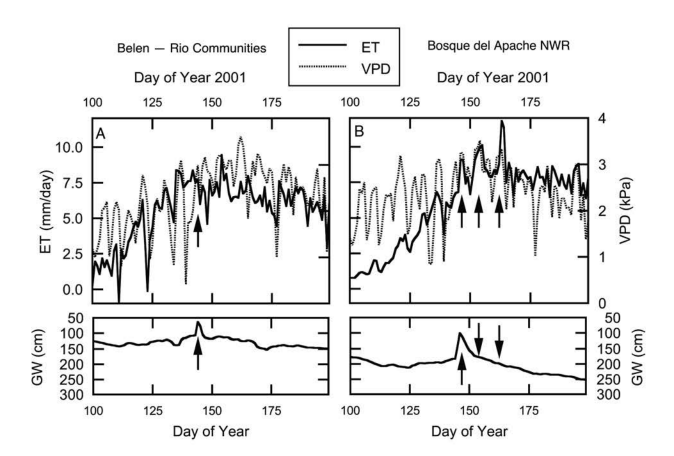


Figure 5:

