A Comparison of Three Methods to Estimate Evapotranspiration in Two Contrasting Loblolly Pine Plantations: Age-Related Changes in Water Use and Drought Sensitivity of Evapotranspiration Components

Jean-Christophe Domec, Ge Sun, Asko Noormets, Michael J. Gavazzi, Emrys A. Treasure, Erika Cohen, Jennifer J. Swenson, Steve G. McNulty, and John S. King

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Keywords: eddy covariance, loblolly pine, Pinus taeda, sapflow, soil moisture probes, water table

Loblolly Pine (Pinus taeda L.) represents one-half of the standing pine volume in the southeastern United States (11.7 million ha) and is by far the single most commercially important forest tree species for the region, with more than 1 billion seedlings planted annually (McKeand et al. 2003). More than 1 million ha of intensively managed loblolly pine plantations are found along the lower coastal plain in eastern North Carolina. Large areas of the North Carolina coastal plain have been drained over the past 300 years, altering more than half of the total forested wetlands of that region (Campbell and Hughes 1991). Unlike upland watersheds dominated by hill slope processes, hydrologic processes on these flat and drained sites are characterized by shallow water tables that are strongly coupled with precipitation and evapotranspiration (ET) (Sun et al. 2002, 2010). For tree species, a shift in ecologic and environmental growing conditions may result in a decline in tree performance and in productivity due to climate change-induced drought stress (Hanson and Weltzin, 2000, Irvine et al. 2004, McDowell et al. 2008). This consideration is especially true for loblolly pine trees that are responsive to prolonged low soil water content (Oren et al. 1998, Ford et al. 2005, Domec et al. 2009), which decreases tree transpiration, gross primary productivity, and net carbon exchange (Noormets et al. 2010, Sun et al. 2010). Soil water content is the essential state variable in hydrologic studies of...
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the land surface because it reflects short-term differences between inputs (e.g., precipitation) and outputs (e.g., ET, runoff, and drainage) of an ecosystem. Most ecohydrologic models are centered on soil water content (e.g., Sperry et al. 1998, Lai and Katul 2000, Barnard et al. 2010). Therefore, accurately defining the relationships between soil water content and water fluxes from the soil and the vegetation is fundamental to improving hydrologic modeling. However, to date the majority of field investigations have relied on monitoring the first 10–40 cm soil profile to characterize whole stand soil water depletion and water use, whereas the rooting zone is usually much deeper (Warren et al. 2005, Miller et al. 2007, Domec et al. 2010).

Variations in water balance across space and time are a function of interactions among species, environmental conditions, stand age, and silvicultural practices (Balodochi et al. 2004, Moore et al. 2004). Evaluating the effects of even-aged forest management on ecosystem water budgets in the coastal regions of the United States has been the focal point of considerable research (McCarthy et al. 1991, Martin 2000, Sun et al. 2000, 2010, Powell et al. 2008) owing to concerns of possible impacts of expansions of plantations on water quality and quality. Widely used hydrologic and water quality models developed for these coastal regions (Amatya and Skaggs 2001) are rarely validated with actual ET, a major hydrologic flux. Furthermore, to evaluate the effects of periodic drought and better predict the impacts of climate change on ecosystem functions from plantations, it is critical to evaluate the response of each of the hydrologic components (Bond et al. 2008, Palmoth et al. 2010). Components of forest water loss may be determined by measuring the simultaneous differences between eddy covariance measurements (ETEC) and tree sapflow (e.g., Oren et al. 1998, Moore et al. 2004, Schwarzel et al. 2009). These two measures can then be compared with soil water content dynamics (Warren et al. 2005, Domec et al. 2010) and with soil evaporation (Raz-Yaseef et al. 2010a). Sap flow usually underestimates tree transpiration because of scaling errors, especially in mixed-species stands (Wilson et al. 2001, Williams et al. 2004, Ford et al. 2007). However, recently it has been shown that in tree plantations with a small number of plant species, ET_EC compared well with sap flow measurements, once corrected for soil evaporation and understory transpiration (Domec et al. 2010, Oishi et al. 2010). This finding suggests that estimation of evaporation from soil and transpiration from the understory might be another source of error in such comparative tests.

Although ET is a key variable that links hydrologic and biologic processes and is critical to modeling watershed hydrology and carbon uptake (Amatya and Skaggs 2001, Sun et al. 2010), large uncertainties remain in its components because it is often too simply derived as the residual of the water balance. In this study, we present a multイヤear comparison of three independent methods that estimate ET and its components. We first compared the ET_EC method with the soil water content variation method (ET_SM). These two techniques give comparable results on shallow soil, but ET_EC is usually larger than ET_SM on deep soils with the differences accentuated during drought (Wilson et al. 2001, Oishi et al. 2008). The observed discrepancies were often attributed to the uncertainty in scaling-up of soil moisture measurements within the entire soil profile (Schwarzel et al. 2009, Oishi et al. 2010), and so we also compared ET_SM estimated from the upper soil profile with ET_SM estimated from the whole soil profile. The third method, water table fluctuation (ET_WT), is based on water level drawdown due to plant uptake and rebound due to underlying gradients in water head (White 1932). Although this technique can be applied on a large scale, it has not been widely used and validated using a comparison with other independent measurements of ET (Vincke and Thiry 2008, Loheide 2008). If proven to compare well, water level fluctuations could provide a useful tool for estimating stand water at low cost over large spatial scales. The objectives of this study were therefore threefold: to characterize the stand water balance in an early- and a mid-rotation loblolly pine plantation using three different approaches; to partition the water use between the loblolly pine trees, the understory trees and soil evaporation; and to evaluate the reliability/representativeness of soil moisture measurements by taking into account the observed variability in soil moisture dynamic by soil depth.

**Methods**

**Sites**

Both study sites are located within the lower coastal plain mixed forest province of North Carolina in the southeastern United States (Noormets et al. 2010, Sun et al. 2010). The loblolly pine plantations are owned and operated by Weyerhaeuser Company. The watersheds are drained with a network of parallel ditches (90–130 cm deep; 90-m spacing) and more widely spaced roadside canals. Drainage lowers the height of the water table, improving site access (management) and tree productivity by reducing stresses caused by excessive soil water conditions during winter months (Kelting et al. 2000). The long-term (1945–2010) average annual precipitation was 1,308 ± 201 mm, evenly distributed throughout the year. Long-term mean annual temperature averaged 15.5° C, with a monthly high temperature occurring in July (26.6° C) and a monthly low occurring in January (6.4° C). The two study sites (US-NC1 and US-NC2 in the Ameriflux database) are 4 km apart, located at 35°11’ N, 76°11’ W and 35°48’ N, 76°40’ W, respectively. The early-rotation plantation (US-NC1) is 48 ha in size, and the mid-rotation stand (US-NC2) is 100 ha. US-NC1 was clearcut in 2004 to remove an 80-year-old native hardwood forest and replanted in 2005 with 1-year-old loblolly pine seedlings (Table 1). The dense understory during the first years was primarily composed of Rubus ursinus (blackberry), Smilax rotundifolia (greenbrier), and Euatorium capillifolium (dog fennel) and reached a height of 2–4 m in 2009. The soil is classified as Cape Fear Series fine, mixed, semiaactive Typic Umbraquult. US-NC2 is a mid-rotation plantation that was established in 1992 after clearcutting the previous mature pine plantation. The histic-mineral soil at this site is classified as Belhaven series. The understory was primarily composed of young red maple (Acer rubrum), devil’s walking stick (Aralia spinosa), pokeweed (Phytolacca americana), beautyberry (Callicarpa
LI-COR Inc., Lincoln, NE). More details on LAI measurements at this site are given in Domec et al. (2009) and Noormets et al. (2010). The optical method was not appropriate because of the open canopy at the early-rotation site, so minimum LAI (winter LAI) was measured destructively on 6–10 trees harvested in late winter every year. The seasonal change in tree LAI at this site was calculated using the seasonal pattern in current-year needle elongation and previous-year needle loss (Domec et al. 2009). The variation curve of total LAI (trees and understory vegetation combined) was determined based on the MODIS-LAI product (Knyazikhin et al. 1998, Oak Ridge National Laboratory Distributed Active Archive Center 2010), which is an 8-day interval time series, and had a total of 43 scenes per year at our sites. Retrieved data were corrected at the early-rotation site for land cover misclassification. MODIS-LAI data were also corrected by applying a canopy extinction coefficient of 0.5 specific to a loblolly pine plantation (Synclair and Knoerr 1982). During winter, when understory LAI was close to 0 (Table 2), MODIS-LAI was highly correlated with tree LAI at both sites (MODIS-LAI = 1.03 trees LAI; r² = 0.83, P < 0.001, data not shown). After field protocols for forest vegetation sampling (Law et al. 2009), at the mid-rotation site we also measured the seasonal change in leaf loss using 30 litter traps (0.18 m² screen-lined laundry baskets) collected every 6 weeks during spring and summer and every 2 weeks during fall and early winter. Litter basket measurements indicated that broadleaf species and understory plants accounted for 16–18% of annual leaf dry biomass. Because specific leaf area of these broadleaf species is approximately 3.5 times higher than pine specific leaf area (Domec et al. 2010), we estimated that understory LAI represents approximately 58% of 1-year pine leaf area (one cohort of needles). Over the 3-year period, one cohort of pine needles taken as winter tree LAI equaled 2.9, putting maximum understory LAI estimated from litter baskets at approximately 1.6. This number was within 10% of the difference (1.7) measured between MODIS-LAI and maximum tree LAI, indicating that these two techniques could be

Table 1. Stand characteristics in 2009 for the early-rotation and the mid-rotation loblolly pine plantations.

<table>
<thead>
<tr>
<th>Soil characteristics²</th>
<th>Early-rotation</th>
<th>Mid-rotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–30 cm</td>
<td>Organic matter/ sandy loam</td>
<td>Organic matter/ sandy loam</td>
</tr>
<tr>
<td>30–60 cm</td>
<td>Sandy clay</td>
<td>Sandy clay</td>
</tr>
<tr>
<td>60–90 cm</td>
<td>Sandy clay</td>
<td>Sandy clay</td>
</tr>
<tr>
<td>90–180 cm</td>
<td>Sandy clay</td>
<td>Sandy clay</td>
</tr>
<tr>
<td>Field capacity (m³ m⁻³)</td>
<td>0.48</td>
<td>0.56</td>
</tr>
<tr>
<td>0–60 cm</td>
<td>0.40</td>
<td>0.37</td>
</tr>
<tr>
<td>60–180 cm</td>
<td>0.145</td>
<td>0.16</td>
</tr>
<tr>
<td>0–60 cm</td>
<td>0.09</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Note that the age of the stand is from planting date and the actual tree ages at both sites were 1 year older (because it included the nursery period). Tree density refers to the number of live trees in 2009 and not to the original planting tree density. LAIs represent minimum and maximum values averaged for 2007 to 2009. Canopy height, tree diameter, stand basal area, and LAI were determined from the 13 vegetation survey plots surrounding the eddy flux tower and followed field protocols for forest vegetation sampling (Law et al. 2009).

¹ Data from Diggs (2004), Grace et al. (2006), and Domec et al. (2010).
² Data from Diggs (2004).

Table 2. Monthly winter (averaged between November and February) ET from ETₑₑₑₑ and its partitioning between the evaporative components (I and Eₑₑₑₑ) and the transpirational components (T and understory transpiration) along with LAI of trees and understory plants.

<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td>4-yr early-rotation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ETₑₑₑₑ (mm mo⁻¹)</td>
<td>32 ± 4</td>
<td>33 ± 3</td>
<td>31 ± 6</td>
</tr>
<tr>
<td>I (mm mo⁻¹)</td>
<td>2 ± 1</td>
<td>2 ± 1</td>
<td>3 ± 2</td>
</tr>
<tr>
<td>Eₑₑₑₑ (mm mo⁻¹)</td>
<td>9 ± 3</td>
<td>9 ± 1</td>
<td>12 ± 2</td>
</tr>
<tr>
<td>T (mm mo⁻¹)</td>
<td>NA</td>
<td>14 ± 1</td>
<td>17 ± 2</td>
</tr>
<tr>
<td>Understory transpiration (mm mo⁻¹) = ETₑₑₑₑ - Eₑₑₑₑ - I</td>
<td>NA</td>
<td>6 ± 3</td>
<td>1 ± 2</td>
</tr>
<tr>
<td>Tree LAI</td>
<td>1.0</td>
<td>1.2</td>
<td>1.4</td>
</tr>
<tr>
<td>Understory LAI</td>
<td>0.1</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>17-yr mid-rotation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ETₑₑₑₑ (mm mo⁻¹)</td>
<td>41 ± 5</td>
<td>44 ± 6</td>
<td>56 ± 3</td>
</tr>
<tr>
<td>I (mm mo⁻¹)</td>
<td>8 ± 1</td>
<td>6 ± 1</td>
<td>9 ± 2</td>
</tr>
<tr>
<td>Eₑₑₑₑ (mm mo⁻¹)</td>
<td>6 ± 2</td>
<td>5 ± 1</td>
<td>6 ± 1</td>
</tr>
<tr>
<td>T (mm mo⁻¹)</td>
<td>25 ± 3</td>
<td>33 ± 2</td>
<td>37 ± 5</td>
</tr>
<tr>
<td>Understory transpiration (mm mo⁻¹) = ETₑₑₑₑ - Eₑₑₑₑ - I</td>
<td>2 ± 2</td>
<td>&lt;1 ± 1</td>
<td>3 ± 3</td>
</tr>
<tr>
<td>Tree LAI</td>
<td>2.9</td>
<td>3.0</td>
<td>3.1</td>
</tr>
<tr>
<td>Understory LAI</td>
<td>0.4</td>
<td>0.1</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Data are means ± SD. NA, not applicable.
used to partition tree LAI between trees and understory plants throughout the year.

**Microclimatic Conditions, Soil Moisture, and Stand Water Balance**

At each site, the following micrometeorological parameters were measured above the canopy: relative humidity and air temperature (HMP45AC; Vaisala, Helsinki, Finland), photosynthetic photon flux density (LI-190; LI-COR, Inc.), and gross precipitation (TE-525; Campbell Scientific, Logan, UT). Data were recorded at a 30-minute interval using multiple dataloggers (CR1000 and CR5000 dataloggers; Campbell Scientific).

A simplified closed water balance equation was developed as

\[ P = ET + D + \Delta S_h \] (1)

where \( P \) denotes gross precipitation, \( ET \) is defined as the sum of soil evaporation (\( E_S \)), tree transpiration (\( T \)), understory transpiration, and canopy interception (\( I \)), \( D \) is drainage flowing out of the watershed, and \( \Delta S_h \) represents the change in soil water storage (all in mm year\(^{-1} \)). Drainage was estimated from previous published relationships relating drainage flow rate measured with a weir and water table elevation recorded by the wells (Diggs 2004, Grace et al. 2006).

Continuous water table fluctuations were recorded at 1-hour intervals with a WL40 pressure transducer (Global Water, Port Orange, FL) monitoring well at each site, located less than 15 m from the eddy flux towers (measurement depth = 220 and 130 cm at the mid-rotation and early-rotation site, respectively). \( \Delta S_h \) is the amount of water that is being added to or removed from the unsaturated soil layers and was calculated using volumetric soil water content (\( \theta \)) measured with soil moisture probes consisting of multiple annular capacitance sensors (Sentek Pty. Ltd., Adelaide, Australia), separated vertically by 10 cm or more (Brooks et al. 2002). Except for the very top sensors (5–15 cm), these probes are relatively insensitive to fluctuations in soil temperature and thus remain highly applicable for establishing the magnitude in \( \theta \) and suited for year-round observation of soil moisture conditions (Warren et al. 2005, 2011). Two probes were installed to a depth of 1.4 m with eight independent sensors per probe at the mid-rotation site. Similarly, two probes were installed to a depth of 0.8 m with six independent sensors per probe at the early-rotation site. At both sites these maximum sampled depths encompassed more than 90% of the rooting zone (Domec et al. 2010). Each sensor was calibrated from soil cores taken at each sensor depth. In addition to the Sentek probes, at each site six independent sensors per probe at the early-rotation site. Each sensor was calibrated from soil cores taken at each site. In addition to the Sentek probes, at each site six independent sensors per probe at the early-rotation site. Each sensor was calibrated from soil cores taken at each site.

\( \Delta S_h \) was determined with the Sentek probes to four extra CS616 time domain reflectometers inserted horizontally at four different depths (0–10, 10–20, 20–30, and 30–40 cm). Such comparisons showed no difference in the pattern in soil moisture between the two systems with a 3–4% difference in \( \theta \) (\( P = 0.39, \) Student’s paired \( t \)-test). To compare across sites and to erase the influence of soil texture on \( \theta \), drought intensity was also quantified in the form of relative extractable soil water (dimensionless), as defined by Bréda et al. (2006).

Closure errors (percentage) for annual water balances were computed based on McCarthy et al. (1991):

\[ \% \text{Error} = \frac{\left| \Delta S_{\theta,m} - \Delta S_{\theta,h} \right|}{Q} \times 100 \] (2)

where \( Q \) is the system flux (in mm) expressed as

\[ Q = (P + D + ET + |\Delta S|)/2 \] (3)

\( \Delta S_{\theta,m} \) represents the modeled \( \Delta S_h \) estimated as the residual in Equation 1, and \( \Delta S_h \) was calculated from the measured change in \( \theta \) from using either the whole soil profile or the upper soil layers (first 30 cm). To scale up to the whole rooting zone when the upper soil layers only are used, \( \Delta S \) of the upper soil (\( \Delta S_{\theta,\text{top 30 cm}} \)) was weighted by the root area (\( A_{\text{root}} \)) profile as detailed in Baldocchi et al. (2004):

\[ \Delta S_h = \Delta S_{\theta,\text{top 30 cm}} \times \frac{A_{\text{root}}}{A_{\text{top 30 cm}}} \] (4)

**Evapotranspiration Based on Unsaturated Soil Moisture Depletion and Water Table Fluctuation**

Total \( ET_{SM} \) was calculated as the difference between the maximum and minimum soil water storage measured within a 24-hour period (Brooks et al. 2002, Warren et al. 2007). Intercepted precipitations were added to the calculations of soil water loss to compare \( ET_{SM} \) with \( ET_{EC} \) values:

\[ ET_{SM} = \sum_{i=1}^{n} (\theta_{d,i-1} - \theta_{d,i})z_i + I_d \] (5)

where \( \theta_{d,i} \) is the volumetric soil water content (m\(^3\) m\(^{-3}\)) of layer \( i \) on day \( d \), \( z_i \) is the thickness of soil layer \( i \), \( n \) is the number of soil layers (5 and 8 layers at the early- and mid-rotation sites, respectively), and \( I \) is the canopy interception. Soil water depletion was not estimated on days after rain events. Soil moisture during such days, as well as between adjacent sensors was interpolated linearly. After \( \Delta S_h \), the estimation of \( ET_{SM} \) based on the upper soil layer measurements was determined by weighting soil water depletion of the upper 30 cm by the root area profile.

In addition to the soil water balance method, we also used the water table fluctuation to estimate ET. The method is based on the assumption that a drop in the water table is due to ET and drainage, and a rise in the water table is due to groundwater recharge (White 1932). Recharge is calculated as the product of the change in water level over time and specific yield. This approach is a gross simplification of a complex phenomenon, involving the movement of water to and from the water table (Healy and Cook 2002, Hill and Neary 2007) but has recently been used successfully in forest ecosystems to estimate ET (Loheide 2008, Vincke and Thiry 2008). Recharge by the water table fluctuation method was estimated with the following equation for days with no precipitation (White 1932, Vincke and Thiry 2008):

\[ ET_{W_T} = (\Delta S/t + 24R) \times Y + I_d \] (6)
where ΔS/t is the change in storage for a day of time t found by linear change in water table depth and R is the recovery rate (per hour): the rate of nighttime positive change in water table height (without transpiration). The specific yields (dimensionless) or soil drainable porosity, Ys, for our studied sites were taken from Diggs (2004). For the soil rooting zone, specific yields varied on average from 0.125 at the mid-rotation site to 0.14 at the early-rotation site (Table 1).

Evapotranspiration Estimated Using the Eddy Covariance Method (ETEC)

The eddy covariance towers were located in the middle of the stands and canopy latent heat (\(\lambda E\)) fluxes were measured using an open-path infrared gas analyzer (LI-7500; LI-COR, Inc.) and a three-dimensional sonic anemometer (CSAT3; Campbell Scientific). The data were processed as reported previously (Noormets et al. 2010, Sun et al. 2010). In brief, the 30-minute mean fluxes of \(H_2O\) were calculated as the covariance of vertical wind speed and the concentration of \(H_2O\), representing the total water loss, including soil evaporation and canopy interception. Data were corrected for the warming of the infrared gas analyzer and for fluctuations in air density (Burban et al. 2008), and data quality was judged by atmospheric stability and flux stationarity during periods of well-developed turbulence as reported previously (Noormets et al. 2008). We also screened spurious or incomplete half-hourly data resulting from system malfunction or environmental disturbance. Gaps in 30-minute ET data, which amounted to 7, 14, and 8% for 2007, 2008, and 2009 at the mid-rotation site and to 4, 24, and 12% at the early-rotation site for 2007, 2008, and 2009, respectively, were filled using empirical monthly correlations between observed ET and Food and Agriculture Organization potential evapotranspiration (PET) (Noormets et al. 2010, Sun et al. 2010). Daily ET_{EC} was estimated by summing half-hourly above canopy eddy covariance ecosystem \(\lambda E\) values.

Components of ET: Canopy Interception, Soil Evaporation, Pine Transpiration, and Understory Transpiration

Interception represents evaporation losses during and after rain from the wet leaf surfaces that intercept rainfall. At the mid-rotation site, I was determined from the difference between rainfall measured above the canopy and throughfall measured under the canopy using 10 manual rain gauges with a site visit every 2–3 weeks. Trunk stemflow was not explicitly considered because it represented less than 1% of precipitation at these sites (Sun et al. 2010). For a given species, I varies with forest age, species, and management factors such as spacing and thinning. For conifers it is primarily a function of tree size and thus LAI (Barbier et al. 2009). For the early-rotation site, I was estimated using a relationship between stand basal area and total precipitation determined on similar loblolly pine stands (Stogsdill et al. 1989).

Soil evaporation (E_s) was measured using an 8100 Automated Soil CO₂ Flux System (LI-COR, Inc.) with 8100-101 and 8100-104 automatic chambers. The instrument was initially designed for soil CO₂ efflux measurements but includes a high-precision \(H_2O\) detector. Furthermore, it has been shown that \(H_2O\) concentration increases linearly in the first 45–80 seconds, allowing reliable estimation of \(E_s\) (Raz-Yaseef et al. 2010a, 2010b). Chamber-based soil vapor flux was measured in two soil collars permanently placed at both sites. Measurements were done every 3–5 weeks throughout the entire research period and maximum daily \(E_s\) was calculated at half-hour time intervals between 1,000 and 1,400 hours corresponding with the time of peak in diurnal ET. Multilinear correlations between \(E_s\) and \(\theta\) and between \(E_s\) and vapor pressure deficit were used to estimate \(E_s\) for other periods (Raz-Yaseef et al. 2010a).

Unlike the eddy covariance technique that measures total ET, the sap flow method only measures \(I\) and does not account for \(F\), \(E_s\), and the water transpired by understory plants. We measured sap flux density per unit of conducting xylem area (\(J_s\), g \(m^{-2} s^{-1}\)) in stem xylem of seven trees at both sites. Stem sap flux measurements were made at 0.60 m above the ground at the early-rotation site and at 1.4 m above the ground at four radial positions at the mid-rotation site using 20-mm heat dissipation probes (Granier 1987). Preliminary results showed that there was no significant difference in azimuthal \(J_s\) within trees (\(P = 0.27, F\)-test), probably owing to the homogeneity of pine trunks. Therefore, we inserted all probes on the north-north-west side of the trees (Domec et al. 2009). Thirty-minute averages of temperature difference data were computed and stored in data loggers (CR10; Campbell Scientific). The sensor signal was converted to \(J_s\) according to Granier (1987), and accounted for the effects of nonzero nighttime fluxes on the signal baseline (Oishi et al. 2008). Using tree sapwood area and stand tree density (Table 1), \(J_s\) was scaled and converted to a tree-scale average \(T\) per unit ground area (mm d⁻¹¹). The total sapwood area of the trees equipped with sapflow probes was estimated from the relationship between sapwood area and diameter determined on 22 felled trees and on the diameter of the measured standing trees. Further details on the sap flow method were reported earlier (Domec et al. 2009, 2010). To estimate missing sap flux data, we developed time series models predicting daily \(T\) from climate variables (vapor pressure deficit and photosynthetically active radiation) according to Ford et al. (2005). Missing data resulting from thunderstorms or from probe destruction by bears occurred less than 8% of the time. Because 80% of the missing data occurred on rainy days, the potential flux and the potential impact of the error associated with this prediction were assumed to be low.

Statistical Analysis

Unless otherwise noted in the text the SAS procedure Proc Mixed, which considers the influence of time on the variances of a variable examined (day of year as the repeated measure), was used to test the differences of the means of key observed micrometeorologic and hydrologic
variables between both sites. The significance of least-squares differences among years was calculated with a multiple range test using the Tukey-Kramer test for unbalanced samples.

Results

Seasonal Soil Water Dynamics

The 2007 and 2008 growing seasons experienced an extreme drought, with annual precipitation being 37–41% (350–400 mm) below the 65-year mean (Figure 1). At the beginning of the study in April–May of 2007, the soil was close to field capacity at both sites with the water table level at 30-cm depth and θ of the entire rooting zone ranging from 0.35 to 0.41 m$^3$ m$^{-3}$ (Figure 2A). The water table fell during the subsequent dry period, and summer and fall water tables in 2007 and 2008 were generally much lower than in 2009 (Figure 2A). In June and again between August and November, gradual soil drying occurred. In the mid-rotation stand, the water table level dropped below 1.9 m, and θ fell to 0.11 m$^3$ m$^{-3}$ for the 0–60 cm layer and to 0.19 m$^3$ m$^{-3}$ for the 60–120 cm layer. At this site, field capacity and 100% relative extractable soil water were almost reached again in 2008 for the deeper soil layers when rain events caused a transient partial recovery of θ (data not shown). A series of winter storm events in October and November 2009 (including Hurricane Ida), with a total of 189 mm of precipitation in 4 days, caused water levels at both stands to rise to near the soil surface. Both the groundwater table level and θ indicated that the early-rotation site was wetter than the mid-rotation site with relative extractable water never declining below 20% (Figure 2B). Even in 2007 and 2008, the soil profile at the early-rotation site was periodically saturated during winter and fall or after significant rain events (storms) in the summer. At the mid-rotation site, soil field capacity was not reached again from March 2008 to November 2009, and the relative extractable soil water remained less than 40% during most of this period (Figure 2B).

Components of Stand Evapotranspiration and Water Balance

Over the 3-year period, ecosystem-level evapotranspiration, as estimated with ET$_{EC}$, was 20–30% higher ($P = 0.02$) at the mid-rotation site (943 ± 35 mm year$^{-1}$) than at the early-rotation site (742 ± 74 mm year$^{-1}$). However, significant differences in ET$_{EC}$ occurred between the first 2 dry years and the last wet year. The severe droughts of 2007 and 2008 resulted in a 16–20% and <5% reduction in ET$_{EC}$ compared with 2009 at the early- and mid-rotation sites, respectively. Between sites, ET$_{EC}$ at the early-rotation site during the 2007 and 2008 droughts was 23% lower ($P = 0.016$) than at the mid-rotation site and only 11% lower ($P = 0.055$) in 2009, when soil water was more available (Table 3). The smaller difference in ET$_{EC}$ between stands in 2009 could in part be attributed to increasing LAI at the early-rotation site (Table 1). At both sites the evaporative demand as characterized by PET did not significantly change between years ($P > 0.34$) (Table 3). However, PET was 15% lower at the early-rotation site in 2007 ($P = 0.02$) but not in 2008 and 2009 ($P > 0.31$). The ET$_{EC}$ change was related to precipitation change at the early-rotation site (ET$_{EC}$ = 0.77 × $P$, $r^2 = 0.81$, $P = 0.02$) but not at the mid-rotation site where the dry years did not significantly reduced ET$_{EC}$ ($P = 0.47$). During the dry years, ET$_{EC}$ at the mid-rotation site was 3–12% higher than $P$ and 20% lower during the wet year (or 246 mm year$^{-1}$ less). As a consequence, over the 3 year-period, the water loss through $D$ at the mid-rotation site represented less than 15% of $P$, compared with more than 23% at the early-rotation site (Table 3). Canopy interception was a small but significant contribution to ET, reaching 10% of annual $P$ on average.

Stand-scale soil evaporation (ES) measurements were estimated to be on average 103 ± 9 and 83 ± 6 mm year$^{-1}$ at the early- and mid-rotation sites, respectively (Table 3). Between 2007 and 2009, there was little variation in ES despite variation in rainfall over this period. Soil evaporation represented up to 13 and 25% of ET in May/June at the mid-rotation and early-rotation site, respectively, but only 7% in September/October at both sites. In the early-rotation stand, annual ES decreased from 16 to 11% from 2007 to 2009 ($P = 0.04$).

Evapotranspiration Method Comparison

At the mid-rotation site, ET estimates from soil water depletion (ET$_{SM}$) were calculated using all soil moisture sensors and therefore represented water used for the first 1.4 m in depth where more than 90% of the roots were located (Table 1). The amount of water used in this 1.4 m of soil reached a maximum of 6.1 mm d$^{-1}$, with an average of 4.2 mm d$^{-1}$ through the summer (Figure 3A–C). At the mid-rotation stand, rates of ET$_{SM}$ were consistent on a daily basis with rates of ET$_{EC}$ (Figure 3B and C). However, the daily variations of ET$_{SM}$ as a function of ET$_{EC}$ showed a clockwise hysteresis, reflecting a lagging effect between soil water depletion and stand water loss (Figure 4). At the mid-rotation site, although monthly ET$_{SM}$ across the 3 years was highly correlated with ET$_{EC}$ (Figure 5), ET$_{EC}$ was 12%
lower than \( \text{ET}_{\text{SM}} \) in the wet year (2009). Moreover, the correlation between \( \text{ET}_{\text{EC}} \) and \( \text{ET}_{\text{SM}} \) was highest for the driest year \((r^2 = 0.79, P = 0.02)\) and lowest for the wettest year \((r^2 = 0.53, P = 0.03)\). At the early-rotation site, monthly \( \text{ET}_{\text{SM}} \) across the 3 years was also correlated with \( \text{ET}_{\text{EC}} \) (Figure 5), and the slope between \( \text{ET}_{\text{SM}} \) and \( \text{ET}_{\text{EC}} \) was not significantly different from 1 \((P = 0.39)\).

\( \text{ET}_{\text{WT}} \) resulted in 3-year mean ET estimates of 839 ± 106 mm year\(^{-1}\) at the early-rotation site and 1,116 ± 19 mm year\(^{-1}\) at the mid-rotation site (Table 3). At both sites,

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**Figure 2.** The annual course of daily mean air temperature (\( \text{Ta} \)), vapor pressure deficit (VPD), and the comparison of daily soil moisture dynamic, water table fluctuation (A), and relative extractable water (B) averaged over the entire soil profile between an early-rotation (●) and a mid-rotation pine (■) site from 2007 to 2009, both located within the outer coastal plain of North Carolina.
Table 3. Sum of ET measured from 2007 to 2009 with three different methods: ET_EC, ET_WT, and ET_SM.

<table>
<thead>
<tr>
<th>4-yr early-rotation</th>
<th>2007</th>
<th>2008</th>
<th>2009</th>
</tr>
</thead>
<tbody>
<tr>
<td>Precipitation (mm yr⁻¹) = kg m⁻² soil yr⁻¹</td>
<td>907</td>
<td>882</td>
<td>1,240</td>
</tr>
<tr>
<td>Throughfall (mm yr⁻¹)</td>
<td>848</td>
<td>795</td>
<td>1,153</td>
</tr>
<tr>
<td>Potential evapotranspiration (mm yr⁻¹)</td>
<td>1,024</td>
<td>1,013</td>
<td>1,089</td>
</tr>
<tr>
<td>Evapotranspiration (mm yr⁻¹)</td>
<td>736</td>
<td>637</td>
<td>886</td>
</tr>
<tr>
<td>Interception (mm yr⁻¹)</td>
<td>35</td>
<td>57</td>
<td>101</td>
</tr>
<tr>
<td>Soil evaporation (mm yr⁻¹)</td>
<td>109 ± 14</td>
<td>111 ± 31</td>
<td>95 ± 19</td>
</tr>
<tr>
<td>Tree transpiration (mm yr⁻¹)</td>
<td>NA</td>
<td>192 ± 26</td>
<td>328 ± 49</td>
</tr>
<tr>
<td>Understory transpiration (mm yr⁻¹) = ET_EC - T - E_S</td>
<td>NA</td>
<td>277 ± 43</td>
<td>363 ± 33</td>
</tr>
<tr>
<td>Drainage (mm yr⁻¹)</td>
<td>248</td>
<td>233</td>
<td>213</td>
</tr>
<tr>
<td>Ground water table fluctuation (mm yr⁻¹)</td>
<td>757</td>
<td>667</td>
<td>1,095</td>
</tr>
<tr>
<td>Understory transpiration (mm yr⁻¹) = ET_WT - T - E_S</td>
<td>NA</td>
<td>309 ± 21</td>
<td>569 ± 31</td>
</tr>
<tr>
<td>Soil water depletion (mm yr⁻¹)</td>
<td>659 ± 51</td>
<td>534 ± 57</td>
<td>881 ± 91</td>
</tr>
<tr>
<td>Understory transpiration (mm yr⁻¹) = ET_SM - T - E_S</td>
<td>NA</td>
<td>175 ± 39</td>
<td>357 ± 41</td>
</tr>
<tr>
<td>Modeled soil water storage (ΔS_m, mm yr⁻¹)</td>
<td>-79</td>
<td>8</td>
<td>138</td>
</tr>
<tr>
<td>% Error in soil water storage from upper soil profile</td>
<td>-8</td>
<td>-5</td>
<td>7</td>
</tr>
<tr>
<td>% Error in soil water storage from whole soil profile</td>
<td>-9</td>
<td>-4</td>
<td>-11</td>
</tr>
</tbody>
</table>

17-yr mid-rotation

| Precipitation (mm yr⁻¹) = kg m⁻² soil yr⁻¹ | 892  | 925  | 1,282 |
| Throughfall (mm yr⁻¹) | 826  | 851  | 1,182 |
| Potential evapotranspiration (mm yr⁻¹) | 1,173 | 1,101 | 1,067 |
| Evapotranspiration (mm yr⁻¹) | 1,011 | 927  | 1,001 |
| Interception (mm yr⁻¹) | 66   | 74   | 57   |
| Soil evaporation (mm yr⁻¹) | 89 ± 21 | 79 ± 19 | 82 ± 11 |
| Tree transpiration (mm yr⁻¹) | 644 ± 71 | 648 ± 82 | 777 ± 79 |
| Understory transpiration (mm yr⁻¹) = ET_EC - T - E_S | 213 ± 34 | 116 ± 20 | 77 ± 19 |
| Drainage (mm yr⁻¹) | 255  | 61   | 196  |
| Ground water table fluctuation (mm yr⁻¹) | 318 ± 40 | 333 ± 69 | 181 ± 30 |
| Understory transpiration (mm yr⁻¹) = ET_WT - T - E_S | 1,117 | 1,134 | 1,097 |
| Soil water depletion (mm yr⁻¹) | 980 ± 141 | 1,004 ± 128 | 1,118 ± 101 |
| Understory transpiration (mm yr⁻¹) = ET_SM - T - E_S | 181 ± 27 | 203 ± 46 | 179 ± 41 |
| Modeled soil water storage (ΔS_m, mm yr⁻¹) | -371 | -63  | 78   |
| % Error in soil water storage from whole soil profile | -5  | 8    | 4    |
| % Error in soil water storage from upper soil profile | -20 | 7    | 27   |

ET_EC was decomposed into T, E_S, and understory transpiration. Values of understory transpiration estimated using ET_WT and ET_SM are also given. NA, not applicable.

ΔS_m represents the modeled ΔS_res estimated as the residual in Equation 1. The closure errors (percentage) for annual water balance are given when soil water storage (ΔS_θ) was either calculated from the upper soil profile or from the whole soil profile (Equation 2).

ET_WT was in good agreement with ET_EC on a monthly basis (Figure 5) but not on a daily basis (Figures 3A and C and 4). On a monthly basis, ET_WT predicted ET_EC better at the mid-rotation than at the early-rotation site with a slope varying around 0.85 (Figure 5). At the mid-rotation site, ET_WT overestimated ET_EC by 12–13% (Table 3; Figure 5) and ET_SM by 10–12% (Table 3; correlation between ET_WT and ET_SM not shown).

Comparison of ET and Soil Water Storage Using Whole-Profile and Upper Soil Water Content Variation

At the early-rotation site, the upper soil layer predicted well the whole soil water content dynamic across the 3-year period (Figure 6A). However, at the mid-rotation site, a counterclockwise hysteresis was apparent between θ measured throughout the whole soil profile and the upper soil profile. ET_SM estimated from the upper 30 cm of soil weighted by the root distribution profile underestimated ET_EC by more than 30% (Figure 6B). Similarly, upper soil θ fluctuations provided a poor estimate of T (Figure 6B). Although the overall slopes between monthly T and ET_SM estimated from either the upper 30 cm of soil (Figure 6B) or from the whole soil profile (data not shown) were similar at both sites (P = 0.31), the overall fit decreased by more than 75%.

Significant negative values of measured ΔS_θ were recorded in 2007 and 2008 as a consequence of sustained ET and low P (Figure 7). At the early-rotation site, remaining soil water storage left was only 75 mm by the end of 2008 (Figure 7), which represented fewer than 3 months of T. Annual water budgets indicated that the modeled ΔS_m (estimated as residual from Equation 1) varied from −371 to 78 mm at the mid-rotation stand and from −79 to 138 mm at the early-rotation stand. It can be noted that at the mid-rotation site, 2 consecutive years such as 2007 would have nearly depleted the estimated 780 mm of available water from the entire soil profile where roots are present. At the early-rotation site, monitoring the first 30 cm of soil moisture represented ΔS_θ accurately, as seen by the low closure errors (Table 3). At the mid-rotation site, ΔS_θ calculated using the first 30 cm of the soil profile induced closure errors for the water balances larger than 20% in both the driest and the wettest year (Table 3).
Partitioning the Effect of Soil Evaporation, Tree Transpiration, and Understory Vegetation on the Water Balance

Despite quantitative differences, the daily $T$ values estimated from sap flow and from $ET_{EC}$ were often qualitatively similar over the years (Figure 3C). During winter, the understory deciduous species at both stands were leafless and the overstory pine trees comprised almost 100% of total stand leaf area (Table 2), and the differences between $T$ and $ET_{EC}$ were explained by $I$ and $E_S$ (Table 2), indicating that understory transpiration was negligible ($< 5$ mm month$^{-1}$). However, during the growing season, understory transpiration accounted for a large component of total $ET_{EC}$, averaging 50 and 14% of whole stand water use at the early- and mid-rotation site, respectively. At the mid-rotation site, the understory component declined from 22% in 2007 to 13% in 2008 to 9% in 2009. This sharp decline could be attributed to a decrease in growing season understory LAI from 1.6 in 2007 to 1.4 in 2008 and to 1.2 in 2009. Across years the correlation between understory LAI and understory transpiration was: $\text{transpiration}_{\text{understory}}$ (mm month$^{-1}$) = $17.6 \times \text{LAI}_{\text{understory}} + 11.6$ ($r^2 = 0.77$, $P < 0.001$) at the early-rotation site; $\text{transpiration}_{\text{understory}}$ (mm month$^{-1}$) = $10.4 \times \text{LAI}_{\text{understory}} (r^2 = 0.49$, $P < 0.01$) at the mid-rotation site. The early-rotation site with higher understory LAI relative to tree LAI (Table 1) had lower ET and higher drainage than the mid-rotation site (Table 3).

The use of $ET_{WT}$ to determine the understory component overestimated understory transpiration by more than 16 and 50% at the early- and mid-rotation sites, respectively (Table 3). Understory transpiration calculated using $ET_{SM}$ was underestimated at the early-rotation site by more than 20%. At the mid-rotation site understory transpiration was overestimated by more than 40% in 2008 and 2009 (Table 3).
Discussion

Evapotranspiration: Comparison of Methods

Consistent with an earlier analysis by Sun et al. (2010) for the same sites and by Gholz and Clark (2002) for a similar landscape, ET values were lower at the early-rotation site than at the mid-rotation site regardless of the method used. The differences were amplified when soil water was less available, suggesting that climatic variability could mask the ET differences among stands at different succession stages. In 2007, the difference in ET between sites was also partly explained by the difference in evaporative demand because PET was higher at the mid-rotation site because of higher surface temperature (Sun et al. 2010). However, during the last 2 years, the lower ET values measured at the early-rotation site were not the consequence of differences in PET. Between 2007 and 2009, the evaporative demand at the early-rotation site increased slightly, mirroring the increase in tree LAI and thus affecting surface albedo (Gibbard et al. 2005, Sun et al. 2010).

Soil moisture sensors have drawbacks for calculating ET, because they must be calibrated using soil cores and they integrate only a small soil volume. We acknowledge that the small number of soil moisture probes added some uncertainty in the absolute magnitude of the soil water fluxes. However, monthly ET_SM values were highly correlated with ET_EC (Figure 5), suggesting that the low number of probes did not affect negatively ET_SM in extrapolating results to the same area of the eddy covariance tower. Moreover, comparisons with the time domain reflectometry probes also suggested that we accurately represented soil moisture. Finally, just as with the towers themselves, which are not replicated, the limited spatial representation of soil moisture is offset by the continuous nature of the monitoring. This allowed us to observe changes in system performance through time and in connection to environmental drivers.
The low correlation between ET_{SM} and ET_{EC} for values lower than 2 mm day\(^{-1}\) may be explained by the underestimation of ET_{EC} on wet days because the sonic anemometer and infrared gas analyzer must be dry to function properly. After rain events, the sensors may thus underestimate evaporation of intercepted water as well as ecosystem transpiration (Stoy et al. 2006). Moreover, at the mid-rotation site, the tap root reached 1.9 m (Table 1), so we cannot reject the possibility of significant storage below the depth measured by our system (deepest sensor at 1.4 m). In addition, the hysteresis observed on a daily basis between ET_{SM} and ET_{EC} values could be caused in part by transpiration of water stored in tree stems and branches (Phillips et al. 2003).

At both sites, values of ET_{WT} were also in good agreement with those for ET_{EC} on a monthly basis but not on a daily basis. At the mid-rotation site, ET_{WT} underpredicted ET_{EC} when soil water uptake was high, probably because soil water was also drawn directly from the saturated zone. The water table fluctuation method provided a point value of recharge computed from the water level rise in a well multiplied by the specific yield (i.e., drainable soil porosity) by assuming that a water level rise is caused by recharge entering at the water table (White 1932). Although simple in concept, this method has drawbacks in its application related to accurate estimation of specific yield and to the assumption that the specific yield between soil layers remains constant. Furthermore, for this study, the period from 01:00 to 04:00 was selected for the recovery analysis. However, the choice of this time period is somewhat subjective, and a different period may be better at other sites (Healy and Cook 2002). The optimal period for analysis may even shift and change in duration throughout the growing season within a site. If the plant has the ability to store a significant volume of water and the water status of the plant has been depleted during the day, uptake of water by the roots may continue for several hours after the plant has stopped transpiring (Phillips et al. 2003). Similarly, because hydraulic redistribution from deep roots is a significant process at these sites (Domec et al., 2010), root water uptake from the saturated zone may never be zero. Therefore, high rates of hydraulic redistribution could also explain the difference between ET_{WT} and ET_{SM} (Warren et al. 2007, 2011). For
this reason, the hours immediately after sunset should not be used to determine the recovery rate ($R$ in Equation 6). However, favorable aspects of this method include its simplicity and cost: it can be applied for any shallow well that taps the water table, and an abundance of available water level data exist, allowing spatial extrapolation.

Contributions of Soil Water Storage to ET: Whole Profile Versus Upper Soil Layer

At the mid-rotation site, soil moisture trends revealed the importance of measuring water content at several depths because less than 20% of the water is stored in the top 30 cm of soil. At this site, the discrepancy between $\theta$ measured throughout the whole soil profile and the upper soil profile there was due to the lower soil layers (60–140 cm) contributing a larger amount of water to daily water depletion relative to the upper layers (10–60 cm) when $\theta$ was low (Meiresonne et al. 2003, Warren et al. 2005, Domec et al. 2010). These results suggest that great care should be taken when one is comparing $T$ and $ET_{EC}$ to soil water dynamics measured from the upper soil profile even after correcting for root profile distribution (Miller et al. 2007). However, some of the differences in soil water storage measured throughout the whole soil profile and the upper soil profile are expected to be dampened by water hydraulically redistributed by deep roots (Warren et al. 2007). At the mid-rotation site, it has been recently shown that during the dry period approximately 80 mm of soil water could replenish the upper soil layers though this phenomenon (Domec et al. 2010).

Figure 6. Daily mean of water content the upper soil layers as a function of soil water content of the entire soil profile (A), and monthly sum stand evapotranspiration ($ET_{EC}$) and pine transpiration ($T$) as a function of soil water depletion of the upper soil layers (B). In B, 95% confidence intervals for the regressions are shown.
Great care should also be taken in the study of ecosystem responses to variation in soil $\theta$, such as soil respiration or soil water uptake, from measurements that rely on the upper soil profile only. Field and modeling studies relying on upper soil moisture to represent whole stand soil water dynamics would underestimate soil water depletion significantly during dry years and overestimate it during wet years. The threshold soil water storage corresponding to a relative extractable soil water of 40–50% was reached quickly in 2007 and stayed for more than a full year (Figure 2; Figure 7). This result indicates that the trees were under constant water stress in 2007 and 2008 and responded by closing their stomata to reduce $T$ (Phillips and Oren 2001, Domec et al. 2009). The threshold of 40–50% for relative extractable water, beyond which $T$ is reduced, has been previously reported in a large number of other tree species and soil types (Brêda et al. 2006, Vincke and Thiry 2008, Gonzalez-Benecke and Martin 2010).

**Partitioning the Effect of Soil Evaporation, Tree Transpiration, and Understory Vegetation on the Water Balance**

Canopy $I$ at the mid-rotation site, one of the two components of stand evaporation was low compared with that for other conifer plantations (Loustau et al. 1992), probably because rainfall events were of short duration but of high intensity (Crockford and Richardson 2000), characteristics of the southeastern United States. Our calculations indicated that $E_S$, the other component of stand evaporation that is often disregarded in forested ecosystems, was a major contributor to ET. Few studies estimate or measure $E_S$, and many assume that under a closed-canopy environment and thick forest floor, $E_S$ can be considered to be minimal because of poor coupling with the atmosphere and high resistance to water vapor loss from dry topsoil (Brêda et al. 2006). However, some reports (Benyon and Doody 2004, Oishi et al. 2008, 2010) also suggest that $E_S$ can contribute approximately 10–20% of total ET. The decrease in $E_S$ at the early-rotation site was attributed to an LAI-mediated decrease in soil temperature over the 3-year period, because stand peak LAI at this site increased from 3.3 in 2007 to 4.2 in 2009. In addition to decreasing the exposure of soil surface to incoming radiation, higher LAI also contributes to increased litter production, which may insulate the ground and suppress $E_S$ (Schaap and Bouten 1997, Raz-Yaseef et al. 2010b).

Although the constant heat sap flux approach is often cited as underpredicting $T$ (Wilson et al. 2001, Brêda et al. 2006; Steppe et al. 2010), there was a good quantitative
agreement between $T$ and ET$_{EC}$ during the winter when the overstory pine trees comprised almost 100% of total stand leaf area (Table 2). This was also probably due to the uniformity of the pine trees, reducing between-tree variability, and also because we measured sap flow across almost the entire sapwood depth, thus reducing within-tree uncertainties. Comparable in methodology to the present study, many studies have compared eddy covariance estimates of ET with sap flux scaled estimates of $T$. These studies generally used the difference in latent heat flux and scaled sap flux to partition evaporative fluxes. However, they did not validate sap flux scaled measurements of $T$ per se (Oren et al. 1998, Kurpius et al. 2003, McCulloh et al. 2007). In contrast, most studies assumed that $E_S$ and understory transpiration are negligible due to closed-canopy conditions, therefore assuming that $T$ is directly comparable with ET.

At the early-rotation site, loblolly pine LAI almost doubled between 2007 and 2009 and the growing season LAI of subdominant trees and understory vegetation doubled whole stand LAI (Table 1), explaining why understory transpiration represented up to 58% of total ET over a season (Table 3). Other studies have shown that understory vegetation can represent a considerable fraction of total LAI (Oren et al. 1987, Loustau and Cochard 1991, Baldocchi et al. 2000, Porté et al. 2009) and be the main driver of the increase in whole stand ET and thus carbon exchange after regeneration or until the stand reaches canopy closure (Jarosz et al. 2008, Domec et al. 2010). However, our comparison of methods to determine ET revealed that great care should be taken when using both soil water balance methods to estimate understory transpiration (Table 2).

As already reported for loblolly pine (Phillips and Oren 2001, Domec et al. 2009), $T$ is responsive to a decrease in $\theta$, with a decrease by more than 30% when relative extractable soil water drops to less than 50%. However, although trees from both plantations were sensitive to drought as seen by the lower $T$ in 2007 and 2008, the mid-rotation stand exhibited little variation in annual ET (Table 2). This result has recently been observed in a forest composed of several tree species (Stoy et al. 2006, Oishi et al. 2010). Therefore, it is possible that on the early-rotation site, acclimation to high levels of water stress in 2007 and 2008 produced foliage with a conservative water use strategy. In a young plantation, chronic levels of water stress may affect the rate of water uptake even when water is available, reflecting a carryover effect of water availability on $T$ (Ewers et al. 1999).

Conclusions

The good agreement between the three methods at a monthly scale gives confidence in these approaches for estimating monthly and annual evapotranspiration. The water table fluctuation method does not require elaborate calibration or data processing, and recharge rates are values that are integrated over large areas and thus represent well the tower foot print. This is a distinct advantage relative to point measurement approaches such as the soil moisture probe methods. The choice of ET methods depends on project objectives and financial constraints. The eddy covariance method gives a high-resolution ET estimate for a homogeneous landscape; thus, it has the advantage for examining ecosystem processes when incorporating with simultaneous measurements of other gases (e.g., CO$_2$ and CH$_3$). However, this method is relatively costly compared with the other ET methods examined in this study. The soil water-based approach is less costly and provides a reasonable estimate of ET at a longer time scale (monthly) although it does not apply to daily ET estimates.

The early-rotation site was very sensitive to summer drought as opposed to the mid-rotation site. At the mid-rotation site, soil moisture trends demonstrated the importance of measuring water content at several depths throughout the rooting zone. Combining measurements of sapflow, soil evaporation, and canopy interception provided insights to the response of water use by plants to climatic variability and stand development. Tree transpiration and evapotranspiration were not directly comparable because understory and soil transpiration rates were not negligible. Pine transpiration showed remarkable consistency as the stands developed, regardless of the accompanying increases in LAI and changes in understory composition from early- to mid-rotation. At the early-rotation site, this conservation of water use was achieved through relatively low transpiration rates (compared with PET and ET) and compensatory transpiration among the overstory trees and understory vegetation. At this site, reduced water availability will affect trees more than understory or soil, whereas at the mid-rotation site, it would affect more understory transpiration, which might be conducive for management efforts.

Literature Cited

BRÉDA, N., R. HUC, A. GRANIER, AND E. DREYER. 2006. Temperate forest trees and stands under severe drought: A review of


the land surface because it reflects short-term differences between inputs (e.g., precipitation) and outputs (e.g., ET, runoff, and drainage) of an ecosystem. Most ecohydrologic models are centered on soil water content (e.g., Sperry et al. 1998, Lai and Katul 2000, Barnard et al. 2010). Therefore, accurately defining the relationships between soil water content and water fluxes from the soil and the vegetation is fundamental to improving hydrologic modeling. However, to date the majority of field investigations have relied on monitoring the first 10–40 cm soil profile to characterize whole stand soil water depletion and water use, whereas the rooting zone is usually much deeper (Warren et al. 2005, Miller et al. 2007, Domec et al. 2010).

Variations in water balance across space and time are a function of interactions among species, environmental conditions, stand age, and silvicultural practices (Baldocchi et al. 2004, Moore et al. 2004). Evaluating the effects of even-aged forest management on ecosystem water budgets in the coastal regions of the United States has been the focal point of considerable research (McCarthy et al. 1991, Martin 2000, Sun et al. 2000, 2010, Powell et al. 2008) owing to concerns of possible impacts of expansions of plantations on water quality and quality. Widely used hydrologic and water quality models developed for these coastal regions (Amatya and Skaggs 2001) are rarely validated with actual ET, a major hydrologic flux. Furthermore, to evaluate the effects of periodic drought and better predict the impacts of climate change on ecosystem functions from plantations, it is critical to evaluate the response of each of the hydrologic components (Bond et al. 2008, Palmroth et al. 2010). Components of forest water loss may be determined by measuring the simultaneous differences between eddy covariance measurements (ETEC) and tree sapflow (e.g., Oren et al. 1998, Moore et al. 2004, Schwarzel et al. 2009). These two measures can then be compared with soil water content dynamics (Warren et al. 2005, Domec et al. 2010) and with soil evaporation (Raz-Yaseef et al. 2010a). Sap flow usually underestimates tree transpiration because of scaling errors, especially in mixed-species stands (Wilson et al. 2001, Williams et al. 2004, Ford et al. 2007). However, recently it has been shown that in tree plantations with a small number of plant species, ET_{EC} compared well with sap flow measurements, once corrected for soil evaporation and understory transpiration (Domec et al. 2010, Oishi et al. 2010). This finding suggests that estimation of evaporation from soil and transpiration from the understory might be another source of error in such comparative tests.

Although ET is a key variable that links hydrologic and biologic processes and is critical to modeling watershed hydrology and carbon uptake (Amatya and Skaggs 2001, Sun et al. 2010), large uncertainties remain in its components because it is often too simply derived as the residual of the water balance. In this study, we present a multyear comparison of three independent methods that estimate ET and its components. We first compared the ET_{EC} method with the soil water content variation method (ET_{SM}). These two techniques give comparable results on shallow soil, but ET_{EC} is usually larger than ET_{SM} on deep soils with the differences accentuated during drought (Wilson et al. 2001, Oishi et al. 2008). The observed discrepancies were often attributed to the uncertainty in scaling-up of soil moisture measurements within the entire soil profile (Schwarzel et al. 2009, Oishi et al. 2010), and so we also compared ET_{SM} estimated from the upper soil profile with ET_{SM} estimated from the whole soil profile. The third method, water table fluctuation (ET_{WT}), is based on water level drawdown due to plant uptake and rebound due to underlying gradients in water head (White 1932). Although this technique can be applied on a large scale, it has not been widely used and validated using a comparison with other independent measurements of ET (Vincke and Thiry 2008, Loheide 2008). If proven to compare well, water level fluctuations could provide a useful tool for estimating stand water at low cost over large spatial scales. The objectives of this study were therefore threefold: to characterize the stand water balance in an early- and a mid-rotation loblolly pine plantation using three different approaches; to partition the water use between the loblolly pine trees, the understory trees and soil evaporation; and to evaluate the reliability/representativeness of soil moisture measurements by taking into account the observed variability in soil moisture dynamic by soil depth.

**Methods**

**Sites**

Both study sites are located within the lower coastal plain mixed forest province of North Carolina in the southeastern United States (Noormets et al. 2010, Sun et al. 2010). The loblolly pine plantations are owned and operated by Weyerhaeuser Company. The watersheds are drained with a network of parallel ditches (90–130 cm deep; 90-m spacing) and more widely spaced roadside canals. Drainage lowers the height of the water table, improving site access (management) and tree productivity by reducing stresses caused by excessive soil water conditions during winter months (Kelting et al. 2000). The long-term (1945–2010) average annual precipitation was 1,308 ± 201 mm, evenly distributed throughout the year. Long-term mean annual temperature averaged 15.5° C, with a monthly high temperature occurring in July (26.6° C) and a monthly low occurring in January (6.4° C). The two study sites (US-NC1 and US-NC2 in the Ameriflux database) are 4 km apart, located at 35°11’ N, 76°11’ W and 35°48’ N, 76°40’ W, respectively. The early-rotation plantation (US-NC1) is 48 ha in size, and the mid-rotation stand (US-NC2) is 100 ha. US-NC1 was clearcut in 2004 to remove an 80-year-old native hardwood forest and replanted in 2005 with 1-year-old loblolly pine seedlings (Table 1). The dense understory during the first years was primarily composed of Related plant species, ETEC compared well with sap flow measurements within the entire soil profile (Schwarzel et al. 2009, Oishi et al. 2010).
Table 1. Stand characteristics in 2009 for the early-rotation and the mid-rotation loblolly pine plantations.

<table>
<thead>
<tr>
<th></th>
<th>Early-rotation stand</th>
<th>Mid-rotation stand</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stand age</td>
<td>5</td>
<td>19</td>
</tr>
<tr>
<td>Canopy height (m)</td>
<td>3.5</td>
<td>18.7</td>
</tr>
<tr>
<td>Tree spacing (m)</td>
<td>1.5 by 6</td>
<td>2.5 by 6</td>
</tr>
<tr>
<td>Density (tree ha(^{-1}))</td>
<td>1,040</td>
<td>635</td>
</tr>
<tr>
<td>Tree dbh (m)</td>
<td>0.14</td>
<td>0.35</td>
</tr>
<tr>
<td>Stand basal area (m(^2) ha(^{-1}))</td>
<td>14.5</td>
<td>56.2</td>
</tr>
<tr>
<td>Tree LAI</td>
<td>1.0–2.0</td>
<td>3.0–4.2</td>
</tr>
<tr>
<td>Total LAI</td>
<td>1.2–4.1</td>
<td>3.1–5.4</td>
</tr>
<tr>
<td>Rooting depth (m)</td>
<td>0.9</td>
<td>1.9</td>
</tr>
</tbody>
</table>

Soil characteristics

<table>
<thead>
<tr>
<th></th>
<th>0–30 cm</th>
<th>30–60 cm</th>
<th>60–90 cm</th>
<th>90–180 cm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic matter/sandy loam</td>
<td></td>
<td>Organic matter/sandy loam</td>
<td>Sandy clay</td>
<td>Sandy clay</td>
</tr>
<tr>
<td>0–60 cm</td>
<td>0.48</td>
<td>0.37</td>
<td></td>
<td></td>
</tr>
<tr>
<td>60–180 cm</td>
<td>0.145</td>
<td>0.16</td>
<td>0.09</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Note that the age of the stand is from planting date and the actual tree ages at both sites were 1 year older (because it included the nursery period). Tree density refers to the number of live trees in 2009 and not to the original planting tree density. LAIs represent minimum and maximum values averaged for 2007 to 2009. Canopy height, tree diameter, stand basal area, and LAI were determined from the 13 vegetation survey plots surrounding the eddy flux tower and followed field protocols for forest vegetation sampling (Law et al. 2009).

(1) Data from Diggs (2004), Grace et al. (2006), and Domec et al. (2010).

American, giant cane (Arundinaria macrostachya), and Meadow grass (Poa spp.) (Domec et al. 2010).

Forest projected leaf area index (LAI) at the mid-rotation site was measured using a LAI-2000 Plant Canopy Analyzer (LI-COR Inc., Lincoln, NE). More details on LAI measurements at this site are given in Domec et al. (2009) and Noormets et al. (2010). The optical method was not appropriate because of the open canopy at the early-rotation site, so minimum LAI (winter LAI) was measured destructively on 6–10 trees harvested in late winter every year. The seasonal change in tree LAI at this site was calculated using the seasonal pattern in current-year needle elongation and previous-year needle loss (Domec et al. 2009). The variation curve of total LAI (trees and understory vegetation combined) was determined based on the MODIS-LAI product (Knyazikhin et al. 1998, Oak Ridge National Laboratory Distributed Active Archive Center 2010), which is an 8-day interval time series, and had a total of 43 scenes per year at our sites. Retrieved data were corrected at the early-rotation site for land cover misclassification. MODIS-LAI data were also corrected by applying a canopy extinction coefficient of 0.5 specific to a loblolly pine plantation (Synclair and Knoerr 1982). During winter, when understory LAI was close to 0 (Table 2), MODIS-LAI was highly correlated with tree LAI at both sites (MODIS-LAI = 1.03 trees LAI; \( r^2 = 0.83, P < 0.001 \), data not shown). After field protocols for forest vegetation sampling (Law et al. 2009), at the mid-rotation site we also measured the seasonal change in leaf loss using 30 litter traps (0.18 m\(^2\) screen-lined laundry baskets) collected every 6 weeks during spring and summer and every 2 weeks during fall and early winter. Litter basket measurements indicated that broadleaf species and understory plants accounted for 16–18% of annual leaf dry biomass. Because specific leaf area of these broadleaf species is approximately 3.5 times higher than pine specific leaf area (Domec et al. 2010), we estimated that understory LAI represented approximately 58% of 1-year pine leaf area (one cohort of needles). Over the 3-year period, one cohort of pine needles taken as winter tree LAI equaled 2.9, putting maximum understory LAI estimated from litter baskets at approximately 1.6. This number was within 10% of the difference (1.7) measured between MODIS-LAI and maximum tree LAI, indicating that these two techniques could be.

Table 2. Monthly winter (averaged between November and February) ET from \( E_{\text{TEC}} \) and its partitioning between the evaporative components \( (I \text{ and } E_s) \) and the transpirational components \( (T \text{ and understory transpiration}) \) along with LAI of trees and understory plants.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>4-yr early-rotation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( E_{\text{TEC}} ) (mm mo(^{-1}))</td>
<td>32 ± 4</td>
<td>33 ± 3</td>
<td>31 ± 6</td>
</tr>
<tr>
<td>( I ) (mm mo(^{-1}))</td>
<td>2 ± 1</td>
<td>2 ± 1</td>
<td>3 ± 2</td>
</tr>
<tr>
<td>( E_s ) (mm mo(^{-1}))</td>
<td>9 ± 3</td>
<td>9 ± 1</td>
<td>12 ± 2</td>
</tr>
<tr>
<td>( T ) (mm mo(^{-1}))</td>
<td>NA</td>
<td>14 ± 1</td>
<td>17 ± 2</td>
</tr>
<tr>
<td>Understory transpiration (mm mo(^{-1})) = ( E_{\text{TEC}} - I - E_s )</td>
<td>NA</td>
<td>6 ± 3</td>
<td>1 ± 2</td>
</tr>
<tr>
<td>Tree LAI</td>
<td>1.0</td>
<td>1.2</td>
<td>1.4</td>
</tr>
<tr>
<td>Understory LAI</td>
<td>0.1</td>
<td>0.3</td>
<td>0.2</td>
</tr>
<tr>
<td>17-yr mid-rotation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( E_{\text{TEC}} ) (mm mo(^{-1}))</td>
<td>41 ± 5</td>
<td>44 ± 6</td>
<td>56 ± 3</td>
</tr>
<tr>
<td>( I ) (mm mo(^{-1}))</td>
<td>8 ± 1</td>
<td>6 ± 1</td>
<td>9 ± 2</td>
</tr>
<tr>
<td>( E_s ) (mm mo(^{-1}))</td>
<td>6 ± 2</td>
<td>5 ± 1</td>
<td>6 ± 1</td>
</tr>
<tr>
<td>( T ) (mm mo(^{-1}))</td>
<td>25 ± 3</td>
<td>33 ± 2</td>
<td>37 ± 5</td>
</tr>
<tr>
<td>Understory transpiration (mm mo(^{-1})) = ( E_{\text{TEC}} - I - E_s )</td>
<td>2 ± 2</td>
<td>&lt;1 ± 1</td>
<td>3 ± 3</td>
</tr>
<tr>
<td>Tree LAI</td>
<td>2.9</td>
<td>3.0</td>
<td>3.1</td>
</tr>
<tr>
<td>Understory LAI</td>
<td>0.4</td>
<td>0.1</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Data are means ± SD. NA, not applicable.
used to partition tree LAI between trees and understory plants throughout the year.

**Microclimatic Conditions, Soil Moisture, and Stand Water Balance**

At each site, the following micrometeorologic parameters were measured above the canopy: relative humidity and air temperature (HMP45AC; Vaisala, Helsinki, Finland), photosynthetic photon flux density (LI-190; LI-COR, Inc.), and gross precipitation (TE-525; Campbell Scientific, Logan, UT). Data were recorded at a 30-minute interval using multiple dataloggers (CR1000 and CR5000 dataloggers; Campbell Scientific).

A simplified closed water balance equation was developed as

\[ P = ET + D + \Delta S_p \]  

(1)

where \( P \) denotes gross precipitation, \( ET \) is defined as the sum of soil evaporation (\( E_S \)), tree transpiration (\( T \)), understory transpiration, and canopy interception (\( I \)), \( D \) is drainage flowing out of the watershed, and \( \Delta S_p \) represents the change in soil water storage (all in mm year\(^{-1}\)). Drainage was estimated from previous published relationships relating drainage flow rate measured with a weir and water table elevation recorded by the wells (Diggs 2004, Grace et al. 2006).

Continuous water table fluctuations were recorded at 1-hour intervals with a WL40 pressure transducer (Global Water, Port Orange, FL) monitoring well at each site, located less than 15 m from the eddy flux towers (measurement depth = 220 and 130 cm at the mid-rotation and early-rotation site, respectively). \( \Delta S_p \) is the amount of water that is being added to or removed from the unsaturated soil layers and was calculated using volumetric soil water content (\( \theta \)) measured with soil moisture probes consisting of multiple annular capacitance sensors (Sentek Pty. Ltd., Adelaide, Australia), separated vertically by 10 cm or more (Brooks et al. 2002). Except for the very top sensors (5–15 cm), these probes are relatively insensitive to fluctuations in soil temperature and thus remain highly applicable for establishing the magnitude in \( \theta \) and suited for year-round observation of soil moisture conditions (Warren et al. 2005, 2011). Two probes were installed to a depth of 1.4 m with eight independent sensors per probe at the mid-rotation site. Similarly, two probes were installed to a depth of 0.8 m with six independent sensors per probe at the early-rotation site. At both sites these maximum sampled depths encompassed more than 90% of the rooting zone (Domec et al. 2010). Each sensor was calibrated from soil cores taken at each sensor depth. In addition to the Sentek probes, at each site \( \theta \) was also measured over the top 30 cm of the soil profile using four vertically inserted CS616 time domain reflectometers (Campbell Scientific). Moreover, to address the uncertainty in the \( \theta \) measurements, we also compared \( \theta \) determined with the Sentek probes to four extra CS616 time domain reflectometers inserted horizontally at four different depths (0–10, 10–20, 20–30, and 30–40 cm). Such comparisons showed no difference in the pattern in soil moisture between the two systems with a 3–4% difference in \( \theta \) (\( P = 0.39 \), Student’s paired t-test). To compare across sites and to erase the influence of soil texture on \( \theta \), drought intensity was also quantified in the form of relative extractable soil water (dimensionless), as defined by Bréda et al. (2006).

Closure errors (percentage) for annual water balances were computed based on McCarthy et al. (1991):

\[ \% \text{Error} = \frac{\Delta S_{\theta,m} - \Delta S_p}{Q} \times 100 \]  

(2)

where \( Q \) is the system flux (in mm) expressed as

\[ Q = (P + D + ET + I) \]  

(3)

\( \Delta S_{\theta,m} \) represents the modeled \( \Delta S_p \) estimated as the residual in Equation 1, and \( \Delta S_p \) was calculated from the measured change in \( \theta \) from using either the whole soil profile or the upper soil layers (first 30 cm). To scale up to the whole rooting zone when the upper soil layers only are used, \( \Delta S \) of the upper soil (\( \Delta S_{\theta, \text{top 30 cm}} \)) was weighted by the root area (\( A_{\text{root}} \)) profile as detailed in Baldocchi et al. (2004):

\[ \Delta S_p = \Delta S_{\theta, \text{top 30 cm}} \frac{A_{\text{root}}}{A_{\text{root 30 cm}}} \]  

(4)

**Evapotranspiration Based on Unsaturated Soil Moisture Depletion and Water Table Fluctuation**

Total \( ET_{SM} \) was calculated as the difference between the maximum and minimum soil water storage measured within a 24-hour period (Brooks et al. 2002, Warren et al. 2007). Intercepted precipitations were added to the calculations of soil water loss to compare \( ET_{SM} \) with \( ET_{EC} \) values:

\[ ET_{SM,i} = \sum_{i=1}^{n} (\theta_{i,d-1} - \theta_{i,d})z_i + I_d \]  

(5)

where \( \theta_{i,d} \) is the volumetric soil water content (\( m^3 \) \( m^{-3} \)) of layer \( i \) on day \( d \), \( z_i \) is the thickness of soil layer \( i \), \( n \) is the number of soil layers (5 and 8 layers at the early- and mid-rotation sites, respectively), and \( I \) is the canopy interception. Soil water depletion was not estimated on days after rain events. Soil moisture during such days, as well as between adjacent sensors was interpolated linearly. After \( \Delta S_p \), the estimation of \( ET_{SM} \) based on the upper soil layer measurements was determined by weighting soil water depletion of the upper 30 cm by the root area profile.

In addition to the soil water balance method, we also used the water table fluctuation to estimate ET. The method is based on the assumption that a drop in the water table is due to ET and drainage, and a rise in the water table is due to groundwater recharge (White 1932). Recharge is calculated as the product of the change in water level over time and specific yield. This approach is a gross simplification of a complex phenomenon, involving the movement of water to and from the water table (Healy and Cook 2002, Hill and Neary 2007) but has recently been used successfully in forest ecosystems to estimate ET (Loheide 2008, Vincke and Thiry 2008). Recharge by the water table fluctuation method was estimated with the following equation for days with no precipitation (White 1932, Vincke and Thiry 2008):

\[ ET_{WT} = (\Delta S/t + 24R) \times Y \]  

(6)
where $\Delta S/t$ is the change in storage for a day of time $t$ found by linear change in water table depth and $R$ is the recovery rate (per hour): the rate of nighttime positive change in water table height (without transpiration). The specific yields (dimensionless) or soil drainable porosity, $Y_s$, for our studied sites were taken from Diggs (2004). For the soil rooting zone, specific yields varied on average from 0.125 at the mid-rotation site to 0.14 at the early-rotation site (Table 1).

**Evapotranspiration Estimated Using the Eddy Covariance Method ($ET_{EC}$)**

The eddy covariance towers were located in the middle of the stands and canopy latent heat ($\lambda E$) fluxes were measured using an open-path infrared gas analyzer (LI-7500; LI-COR, Inc.) and a three-dimensional sonic anemometer (CSAT3; Campbell Scientific). The data were processed as reported previously (Noormets et al. 2010, Sun et al. 2010). In brief, the 30-minute mean fluxes of $H_2O$ were calculated as the covariance of vertical wind speed and the concentration of $H_2O$, representing the total water loss, including soil evaporation and canopy interception. Data were corrected for the warming of the infrared gas analyzer and for fluctuations in air density (Burbank et al. 2008), and data quality was judged by atmospheric stability and flux stationarity during periods of well-developed turbulence as reported previously (Noormets et al. 2008). We also screened spurious or incomplete half-hourly data resulting from system malfunction or environmental disturbance. Gaps in 30-minute ET data, which amounted to 7, 14, and 8% for 2007, 2008, and 2009 at the mid-rotation site and to 4, 24, and 12% at the early-rotation site for 2007, 2008, and 2009, respectively, were filled using empirical monthly correlations between observed ET and Food and Agriculture Organization potential evapotranspiration (PET) (Noormets et al. 2010, Sun et al. 2010). Daily $ET_{EC}$ was estimated by summing half-hourly above canopy eddy covariance ecosystem $\lambda E$ values.

**Components of ET: Canopy Interception, Soil Evaporation, Pine Transpiration, and Understory Transpiration**

Interception represents evaporation losses during and after rain from the wet leaf surfaces that intercept rainfall. At the mid-rotation site, $I$ was determined from the difference between rainfall measured above the canopy and throughfall measured under the canopy using 10 manual rain gauges with a site visit every 2–3 weeks. Trunk stemflow was not explicitly considered because it represented less than 1% of precipitation at these sites (Sun et al. 2010). For a given species, $I$ varies with forest age, species, and management factors such as spacing and thinning. For conifers it is primarily a function of tree size and thus LAI (Barbier et al. 2009). For the early-rotation site, $I$ was estimated using a relationship between stand basal area and total precipitation determined on similar loblolly pine stands (Stogsdill et al. 1989).

Soil evaporation ($E_S$) was measured using an 8100 Au-
tomated Soil CO$_2$ Flux System (LI-COR, Inc.) with 8100-101 and 8100-104 automatic chambers. The instrument was initially designed for soil CO$_2$ efflux measurements but includes a high-precision $H_2O$ detector. Furthermore, it has been shown that $H_2O$ concentration increases linearly in the first 45–80 seconds, allowing reliable estimation of $E_S$ (Raz-Yaseef et al. 2010a, 2010b). Chamber-based soil vapor flux was measured in two soil collars permanently placed at both sites. Measurements were done every 3–5 weeks throughout the entire research period and maximum daily $E_S$ was calculated at half-hour time intervals between 1,000 and 1,400 hours corresponding with the time of peak in diurnal ET. Multilinear correlations between $E_S$ and $\theta$ and between $E_S$ and vapor pressure deficit were used to estimate $E_S$ for other periods (Raz-Yaseef et al. 2010a).

Unlike the eddy covariance technique that measures total ET, the sap flow method only measures $T$ and does not account for $I$, $E_S$, and the water transpired by understory plants. We measured sap flux density per unit of conducting xylem area ($J_s$, g m$^{-2}$ s$^{-1}$) in stem xylem of seven trees at both sites. Stem sap flux measurements were made at 0.60 m above the ground at the early-rotation site and at 1.4 m above the ground at four radial positions at the mid-rotation site using 20-mm heat dissipation probes (Granier 1987). Preliminary results showed that there was no significant difference in azimuthal $J_s$ within trees ($P = 0.27$, $F$-test), probably owing to the homogeneity of pine trunks. Therefore, we inserted all probes on the north-north-west side of the trees (Domec et al. 2009). Thirty-minute averages of temperature difference data were computed and stored in data loggers (CR10; Campbell Scientific). The sensor signal was converted to $J_s$, according to Granier (1987), and accounted for the effects of nonzero nighttime fluxes on the signal baseline (Oiishi et al. 2008). Using tree sapwood area and stand tree density (Table 1), $J_s$ was scaled and converted to a tree-scale average $T$ per unit ground area (mm d$^{-1}$). The total sapwood area of the trees equipped with sapflow probes was estimated from the relationship between sapwood area and diameter determined on 22 felled trees and on the diameter of the measured standing trees. Further details on the sap flow method were reported earlier (Domec et al. 2009, 2010). To estimate missing sap flux data, we developed time series models predicting daily $T$ from climate variables (vapor pressure deficit and photosynthetically active radiation) according to Ford et al. (2005). Missing data resulting from thunderstorms or from probe destruction by bears occurred less than 8% of the time. Because 80% of the missing data occurred on rainy days, the potential flux and the potential impact of the error associated with this prediction were assumed to be low.

**Statistical Analysis**

Unless otherwise noted in the text the SAS procedure Proc Mixed, which considers the influence of time on the variances of a variable examined (day of year as the repeated measure), was used to test the differences of the means of key observed micrometeorologic and hydrologic...
variables between both sites. The significance of least-squares differences among years was calculated with a multiple range test using the Tukey-Kramer test for unbalanced samples.

Results

Seasonal Soil Water Dynamics

The 2007 and 2008 growing seasons experienced an extreme drought, with annual precipitation being 37–41% (350–400 mm) below the 65-year mean (Figure 1). At the beginning of the study in April–May of 2007, the soil was close to field capacity at both sites with the water table level at 30-cm depth and $\theta$ of the entire rooting zone ranging from 0.35 to 0.41 m$^3$ m$^{-3}$ (Figure 2A). The water table fell during the subsequent dry period, and summer and fall water tables in 2007 and 2008 were generally much lower than in 2009 (Figure 2A). In June and again between August and November, gradual soil drying occurred. In the mid-rotation stand, the water table level dropped below 1.9 m, and $\theta$ fell to 0.11 m$^3$ m$^{-3}$ for the 0–60 cm layer and to 0.19 m$^3$ m$^{-3}$ for the 60–120 cm layer. At this site, field capacity and 100% relative extractable soil water were almost reached again in 2008 for the deeper soil layers when rain events caused a transient partial recovery of $\theta$ (data not shown). A series of winter storm events in October and November 2009 (including Hurricane Ida), with a total of 189 mm of precipitation in 4 days, caused water levels at both stands to rise to near the soil surface. Both the groundwater table level and $\theta$ indicated that the early-rotation site was wetter than the mid-rotation site with relative extractable water never declining below 20% (Figure 2B). Even in 2007 and 2008, the soil profile at the early-rotation site was periodically saturated during winter and fall or after significant rain events (storms) in the summer. At the mid-rotation site, soil field capacity was not reached again from March 2008 to November 2009, and the relative extractable soil water remained less than 40% during most of this period (Figure 2B).

Components of Stand Evapotranspiration and Water Balance

Over the 3-year period, ecosystem-level evapotranspiration, as estimated with $ET_{EC}$, was 20–30% higher ($P = 0.02$) at the mid-rotation site (943 ± 35 mm year$^{-1}$) than at the early-rotation site (742 ± 74 mm year$^{-1}$). However, significant differences in $ET_{EC}$ occurred between the first 2 dry years and the last wet year. The severe droughts of 2007 and 2008 resulted in a 16–20% and <5% reduction in $ET_{EC}$ compared with 2009 at the early- and mid-rotation sites, respectively. Between sites, $ET_{EC}$ at the early-rotation site during the 2007 and 2008 droughts was 23% lower ($P = 0.016$) than at the mid-rotation site and only 11% lower ($P = 0.055$) in 2009, when soil water was more available (Table 3). The smaller difference in $ET_{EC}$ between stands in 2009 could in part be attributed to increasing LAI at the early-rotation site (Table 1). At both sites the evaporative demand as characterized by PET did not significantly change between years ($P > 0.34$) (Table 3). However, PET was 15% lower at the early-rotation site in 2007 ($P = 0.02$) but not in 2008 and 2009 ($P > 0.31$). The $ET_{EC}$ change was related to precipitation change at the early-rotation site ($ET_{EC} = 0.77 \times P$, $r^2 = 0.81$, $P = 0.02$) but not at the mid-rotation site where the dry years did not significantly reduced $ET_{EC}$ ($P = 0.47$). During the dry years, $ET_{EC}$ at the mid-rotation site was 3–12% higher than $P$ and 20% lower during the wet year (or 246 mm year$^{-1}$ less). As a consequence, over the 3 year-period, the water loss through $D$ at the mid-rotation site represented less than 15% of $P$, compared with more than 23% at the early-rotation site (Table 3). Canopy interception was a small but significant contribution to ET, reaching 10% of annual $P$ on average.

Stand-scale soil evaporation ($E_s$) measurements were estimated to be on average 103 ± 9 and 83 ± 6 mm year$^{-1}$ at the early- and mid-rotation sites, respectively (Table 3). Between 2007 and 2009, there was little variation in $E_s$ despite variation in rainfall over this period. Soil evaporation represented up to 13 and 25% of ET in May/June at the mid-rotation and early-rotation site, respectively, but only 7% in September/October at both sites. In the early-rotation stand, annual $E_s$ decreased from 16 to 11% from 2007 to 2009 ($P = 0.04$).

Evapotranspiration Method Comparison

At the mid-rotation site, ET estimates from soil water depletion ($ET_{SM}$) were calculated using all soil moisture sensors and therefore represented water used for the first 1.4 m in depth where more than 90% of the roots were located (Table 1). The amount of water used in this 1.4 m of soil reached a maximum of 6.1 mm d$^{-1}$, with an average of 4.2 mm d$^{-1}$ through the summer (Figure 3A–C). At the mid-rotation stand, rates of $ET_{SM}$ were consistent on a daily basis with rates of $ET_{EC}$ (Figure 3B and C). However, the daily variations of $ET_{SM}$ as a function of $ET_{EC}$ showed a clockwise hysteresis, reflecting a lagging effect between soil water depletion and stand water loss (Figure 4). At the mid-rotation site, although monthly $ET_{SM}$ across the 3 years was highly correlated with $ET_{EC}$ (Figure 5), $ET_{EC}$ was 12%
lower than ET$_{SM}$ in the wet year (2009). Moreover, the correlation between ET$_{EC}$ and ET$_{SM}$ was highest for the driest year ($r^2 = 0.79$, $P = 0.02$) and lowest for the wettest year ($r^2 = 0.53$, $P = 0.03$). At the early-rotation site, monthly ET$_{SM}$ across the 3 years was also correlated with ET$_{EC}$ (Figure 5), and the slope between ET$_{SM}$ and ET$_{EC}$ was not significantly different from 1 ($P = 0.39$).

ET$_{WT}$ resulted in 3-year mean ET estimates of 839 ± 106 mm year$^{-1}$ at the early-rotation site and 1,116 ± 19 mm year$^{-1}$ at the mid-rotation site (Table 3). At both sites,
Table 3. Sum of ET measured from 2007 to 2009 with three different methods: ET$_{EC}$, ET$_{WT}$, and ET$_{SM}$.

<table>
<thead>
<tr>
<th></th>
<th>2007</th>
<th>2008</th>
<th>2009</th>
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<tbody>
<tr>
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<tr>
<td>Precipitation (mm yr$^{-1}$)</td>
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<tr>
<td>Interception (mm yr$^{-1}$)</td>
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<td>57</td>
<td>101</td>
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<tr>
<td>Soil evaporation (mm yr$^{-1}$)</td>
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<td>363 + 33</td>
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<td>213</td>
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<td>Ground water table fluctuation (mm yr$^{-1}$)</td>
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<tr>
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<td>Ground water table fluctuation (mm yr$^{-1}$)</td>
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<td>333 + 69</td>
<td>181 + 30</td>
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<td>Soil water depletion (mm yr$^{-1}$)</td>
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<td>1,004 + 128</td>
<td>1,118 + 101</td>
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<td>Understory transpiration (mm yr$^{-1}$)</td>
<td>181 + 27</td>
<td>203 + 46</td>
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<td>Modeled soil water storage ($\Delta S_{soil,mm}$ mm yr$^{-1}$)</td>
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<td>% Error in soil water storage from upper soil profilea</td>
<td>-20</td>
<td>7</td>
<td>27</td>
</tr>
</tbody>
</table>

ET$_{EC}$ was decomposed into $I$, $E_s$, $T$, and understory transpiration. Values of understory transpiration estimated using ET$_{WT}$ and ET$_{SM}$ are also given. NA, not applicable.

$^a$$\Delta S_{soil}$ represents the modeled $\Delta S_{\theta}$ estimated as the residual in Equation 1. The closure errors (percentage) for annual water balance are given when soil water storage ($\Delta S_{\theta}$) was either calculated from the upper soil profile or from the whole soil profile (Equation 2).

ET$_{WT}$ was in good agreement with ET$_{EC}$ on a monthly basis (Figure 5) but not on a daily basis (Figures 3A and C and 4). On a monthly basis, ET$_{WT}$ predicted ET$_{EC}$ better at the mid-rotation than at the early-rotation site with a slope varying around 0.85 (Figure 5). At the mid-rotation site, ET$_{WT}$ overestimated ET$_{EC}$ by 12–13% (Table 3; Figure 5) and ET$_{SM}$ by 10–12% (Table 3; correlation between ET$_{WT}$ and ET$_{SM}$ not shown).

**Comparison of ET and Soil Water Storage Using Whole-Profile and Upper Soil Water Content Variation**

At the early-rotation site, the upper soil layer predicted well the whole soil water content dynamic across the 3-year period (Figure 6A). However, at the mid-rotation site, a counterclockwise hysteresis was apparent between $\theta$ measured throughout the whole soil profile and the upper soil profile. ET$_{SM}$ estimated from the upper 30 cm of soil weighted by the root distribution profile underestimated ET$_{EC}$ by more than 30% (Figure 6B). Similarly, upper soil $\theta$ fluctuations provided a poor estimate of $T$ (Figure 6B). Although the overall slopes between monthly $T$ and ET$_{SM}$ estimated from either the upper 30 cm of soil (Figure 6B) or from the whole soil profile (data not shown) were similar at both sites ($P = 0.31$), the overall fit decreased by more than 75%.

Significant negative values of measured $\Delta S_{\theta}$ were recorded in 2007 and 2008 as a consequence of sustained ET and low $P$ (Figure 7). At the early-rotation site, remaining soil water storage left was only 75 mm by the end of 2008 (Figure 7), which represented fewer than 3 months of $T$. Annual water budgets indicated that the modeled $\Delta S_{soil}$ (estimated as residual from Equation 1) varied from −731 to 77 mm at the mid-rotation stand and from −79 to 138 mm at the early-rotation stand. It can be noted that at the mid-rotation site, 2 consecutive years such as 2007 would have nearly depleted the estimated 780 mm of available water from the entire soil profile where roots are present. At the early-rotation site, monitoring the first 30 cm of soil moisture represented $\Delta S_{\theta}$ accurately, as seen by the low closure errors (Table 3). At the mid-rotation site, $\Delta S_{\theta}$ calculated using the first 30 cm of the soil profile induced closure errors for the water balances larger than 20% in both the driest and the wettest year (Table 3).
Partitioning the Effect of Soil Evaporation, Tree Transpiration, and Understory Vegetation on the Water Balance

Despite quantitative differences, the daily $T$ values estimated from sap flow and from $ET_{EC}$ were often qualitatively similar over the years (Figure 3C). During winter, the understory deciduous species at both stands were leafless and the overstory pine trees comprised almost 100% of total stand leaf area (Table 2), and the differences between $T$ and $ET_{EC}$ were explained by $I$ and $E_S$ (Table 2), indicating that understory transpiration was negligible (<5 mm month$^{-1}$). However, during the growing season, understory transpiration accounted for a large component of total $ET_{EC}$, averaging 50 and 14% of whole stand water use at the early- and mid-rotation site, respectively. At the mid-rotation site, the understory component declined from 22% in 2007 to 13% in 2008 to 9% in 2009. This sharp decline could be attributed to a decrease in growing season understory LAI from 1.6 in 2007 to 1.4 in 2008 and to 1.2 in 2009. Across years the correlation between understory LAI and understory transpiration was: $\text{transpiration}_{\text{understory}}$ (mm month$^{-1}$) = 17.6 $\times$ LAI$_{\text{understory}}$ + 11.6 ($r^2 = 0.77$, $P < 0.001$) at the early-rotation site; $\text{transpiration}_{\text{understory}}$ (mm month$^{-1}$) = 10.4 $\times$ LAI$_{\text{understory}}$ ($r^2 = 0.49$, $P < 0.01$) at the mid-rotation site. The early-rotation site with higher understory LAI relative to tree LAI (Table 1) had lower ET and higher drainage than the mid-rotation site (Table 3).

The use of ET$_{WT}$ to determine the understory component overestimated understory transpiration by more than 16 and 50% at the early- and mid-rotation sites, respectively (Table 3). Understory transpiration calculated using ET$_{SM}$ was underestimated at the early-rotation site by more than 20%. At the mid-rotation site understory transpiration was overestimated by more than 40% in 2008 and 2009 (Table 3).
Discussion

Evapotranspiration: Comparison of Methods

Consistent with an earlier analysis by Sun et al. (2010) for the same sites and by Gholz and Clark (2002) for a similar landscape, ET values were lower at the early-rotation site than at the mid-rotation site regardless of the method used. The differences were amplified when soil water was less available, suggesting that climatic variability could mask the ET differences among stands at different succession stages. In 2007, the difference in ET between sites was also partly explained by the difference in evaporative demand because PET was higher at the mid-rotation site because of higher surface temperature (Sun et al. 2010). However, during the last 2 years, the lower ET values measured at the early-rotation site were not the consequence of differences in PET. Between 2007 and 2009, the evaporative demand at the early-rotation site increased slightly, mirroring the increase in tree LAI and thus affecting surface albedo (Gibbard et al. 2005, Sun et al. 2010).

Soil moisture sensors have drawbacks for calculating ET, because they must be calibrated using soil cores and they integrate only a small soil volume. We acknowledge that the small number of soil moisture probes added some uncertainty in the absolute magnitude of the soil water fluxes. However, monthly ET$_{SM}$ values were highly correlated with ET$_{EC}$ (Figure 5), suggesting that the low number of probes did not affect negatively ET$_{SM}$ in extrapolating results to the same area of the eddy covariance tower. Moreover, comparisons with the time domain reflectometry probes also suggested that we accurately represented soil moisture. Finally, just as with the towers themselves, which are not replicated, the limited spatial representation of soil moisture is offset by the continuous nature of the monitoring. This allowed us to observe changes in system performance through time and in connection to environmental drivers.
The low correlation between ET<sub>SM</sub> and ET<sub>EC</sub> for values lower than 2 mm day<sup>−1</sup> may be explained by the underestimation of ET<sub>EC</sub> on wet days because the sonic anemometer and infrared gas analyzer must be dry to function properly. After rain events, the sensors may thus underestimate evaporation of intercepted water as well as ecosystem transpiration (Stoy et al. 2006). Moreover, at the mid-rotation site, the tap root reached 1.9 m (Table 1), so we cannot reject the possibility of significant storage below the depth measured by our system (deepest sensor at 1.4 m). In addition, the hysteresis observed on a daily basis between ET<sub>SM</sub> and ET<sub>EC</sub> values could be caused in part by transpiration of water stored in tree stems and branches (Phillips et al. 2003).

At both sites, values of ET<sub>WT</sub> were also in good agreement with those for ET<sub>EC</sub> on a monthly basis but not on a daily basis. At the mid-rotation site, ET<sub>WT</sub> underpredicted ET<sub>EC</sub> when soil water uptake was high, probably because soil water was also drawn directly from the saturated zone. The water table fluctuation method provided a point value of recharge computed from the water level rise in a well multiplied by the specific yield (i.e., drainable soil porosity) by assuming that a water level rise is caused by recharge entering at the water table (White 1932). Although simple in concept, this method has drawbacks in its application related to accurate estimation of specific yield and to the assumption that the specific yield between soil layers remains constant. Furthermore, for this study, the period from 01:00 to 04:00 was selected for the recovery analysis. However, the choice of this time period is somewhat subjective, and a different period may be better at other sites (Healy and Cook 2002). The optimal period for analysis may even shift and change in duration throughout the growing season within a site. If the plant has the ability to store a significant volume of water and the water status of the plant has been depleted during the day, uptake of water by the roots may continue for several hours after the plant has stopped transpiring (Phillips et al. 2003). Similarly, because hydraulic redistribution from deep roots is a significant process at these sites (Domec et al., 2010), root water uptake from the saturated zone may never be zero. Therefore, high rates of hydraulic redistribution could also explain the difference between ET<sub>WT</sub> and ET<sub>SM</sub> (Warren et al. 2007, 2011). For

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Figure 5. Monthly sum of water uptake in the water table (ET<sub>WT</sub>), soil water depletion of the entire soil profile (ET<sub>SM</sub>), and pine transpiration (T) as a function of stand evapotranspiration (ET<sub>EC</sub>) (95% confidence intervals for the regressions are shown).
this reason, the hours immediately after sunset should not be used to determine the recovery rate ($R$ in Equation 6). However, favorable aspects of this method include its simplicity and cost: it can be applied for any shallow well that taps the water table, and an abundance of available water level data exist, allowing spatial extrapolation.

**Contributions of Soil Water Storage to ET: Whole Profile Versus Upper Soil Layer**

At the mid-rotation site, soil moisture trends revealed the importance of measuring water content at several depths because less than 20% of the water is stored in the top 30 cm of soil. At this site, the discrepancy between $\theta$ measured throughout the whole soil profile and the upper soil profile there was due to the lower soil layers (60–140 cm) contributing a larger amount of water to daily water depletion relative to the upper layers (10–60 cm) when $\theta$ was low (Meiresonne et al. 2003, Warren et al. 2005, Domec et al. 2010). These results suggest that great care should be taken when one is comparing $T$ and $\text{ET}_{\text{EC}}$ to soil water dynamics measured from the upper soil profile even after correcting for root profile distribution (Miller et al. 2007). However, some of the differences in soil water storage measured throughout the whole soil profile and the upper soil profile are expected to be dampened by water hydraulically redistributed by deep roots (Warren et al. 2007). At the mid-rotation site, it has been recently shown that during the dry period approximately 80 mm of soil water could replenish the upper soil layers though this phenomenon (Domec et al. 2010).

Figure 6. Daily mean of water content the upper soil layers as a function of soil water content of the entire soil profile (A), and monthly sum stand evapotranspiration ($\text{ET}_{\text{EC}}$) and pine transpiration ($T$) as a function of soil water depletion of the upper soil layers (B). In B, 95% confidence intervals for the regressions are shown.
Great care should also be taken in the study of ecosystem responses to variation in soil $\theta$, such as soil respiration or soil water uptake, from measurements that rely on the upper soil profile only. Field and modeling studies relying on upper soil moisture to represent whole stand soil water dynamics would underestimate soil water depletion significantly during dry years and overestimate it during wet years. The threshold soil water storage corresponding to a relative extractable soil water of 40–50% was reached quickly in 2007 and stayed for more than a full year (Figure 2; Figure 7). This result indicates that the trees were under constant water stress in 2007 and 2008 and responded by closing their stomata to reduce $T$ (Phillips and Oren 2001, Domec et al. 2009). The threshold of 40–50% for relative extractable water, beyond which $T$ is reduced, has been previously reported in a large number of other tree species and soil types (Brêda et al. 2006, Vincke and Thiry 2008, Gonzalez-Benecke and Martin 2010).

**Partitioning the Effect of Soil Evaporation, Tree Transpiration, and Understory Vegetation on the Water Balance**

Canopy $I$ at the mid-rotation site, one of the two components of stand evaporation was low compared with that for other conifer plantations (Loustau et al. 1992), probably because rainfall events were of short duration but of high intensity (Crockford and Richardson 2000), characteristics of the southeastern United States. Our calculations indicated that $E_S$, the other component of stand evaporation that is often disregarded in forested ecosystems, was a major contributor to ET. Few studies estimate or measure $E_S$, and many assume that under a closed-canopy environment and thick forest floor, $E_S$ can be considered to be minimal because of poor coupling with the atmosphere and high resistance to water vapor loss from dry topsoil (Brêda et al. 2006). However, some reports (Benyon and Doody 2004, Oishi et al. 2008, 2010) also suggest that $E_S$ can contribute approximately 10–20% of total ET. The decrease in $E_S$ at the early-rotation site was attributed to an LAI-mediated decrease in soil temperature over the 3-year period, because stand peak LAI at this site increased from 3.3 in 2007 to 4.2 in 2009. In addition to decreasing the exposure of soil surface to incoming radiation, higher LAI also contributes to increased litter production, which may insulate the ground and suppress $E_S$ (Schaap and Bouten 1997, Raz-Yaseef et al. 2010b).

Although the constant heat sap flux approach is often cited as underpredicting $T$ (Wilson et al. 2001, Brêda et al. 2006; Steppe et al. 2010), there was a good quantitative...
agreement between $T$ and $ET_{rc}$ during the winter when the overstory pine trees comprised almost 100% of total stand leaf area (Table 2). This was also probably due to the uniformity of the pine trees, reducing between-tree variability, and also because we measured sap flow across almost the entire sapwood depth, thus reducing within-tree uncertainties. Comparable in methodology to the present study, many studies have compared eddy covariance estimates of ET with sap flux scaled estimates of $T$. These studies generally used the difference in latent heat flux and scaled sap flux to partition evaporative fluxes. However, they did not validate sap flux scaled measurements of $T$ per se (Oren et al. 1998, Kurpius et al. 2003, McCulloh et al. 2007). In contrast, most studies assumed that $E_s$ and understory transpiration are negligible due to closed-canopy conditions, therefore assuming that $T$ is directly comparable with ET.

At the early-rotation site, loblolly pine LAI almost doubled between 2007 and 2009 and the growing season LAI of subdominant trees and understory vegetation doubled whole stand LAI (Table 1), explaining why understory transpiration represented up to 58% of total ET over a season (Table 3). Other studies have shown that understory vegetation can represent a considerable fraction of total LAI (Oren et al. 1987, Loustau and Cochard 1991, Baldocchi et al. 2000, Porté et al. 2009) and be the main driver of the increase in whole stand ET and thus carbon exchange after regeneration or until the stand reaches canopy closure (Jarosz et al. 2008, Domec et al. 2010). However, our comparison of methods to determine ET revealed that great care should be taken when using both soil water balance methods to estimate understory transpiration (Table 2).

As already reported for loblolly pine (Phillips and Oren 2001, Domec et al. 2009), $T$ is responsive to a decrease in $\theta$, with a decrease by more than 30% when relative extractable soil water drops to less than 50%. However, although trees from both plantations were sensitive to drought as seen by the lower $T$ in 2007 and 2008, the mid-rotation stand exhibited little variation in annual ET (Table 2). This result has recently been observed in a forest composed of several tree species (Stoy et al. 2006, Oishi et al. 2010). Therefore, it is possible that on the early-rotation site, acclimation to high levels of water stress in 2007 and 2008 produced foliage with a conservative water use strategy. In a young plantation, chronic levels of water stress may affect the rate of water uptake even when water is available, reflecting a carryover effect of water availability on $T$ (Ewers et al. 1999).

**Conclusions**

The good agreement between the three methods at a monthly scale gives confidence in these approaches for estimating monthly and annual evapotranspiration. The water table fluctuation method does not require elaborate calibration or data processing, and recharge rates are values that are integrated over large areas and thus represent well the tower footprint. This is a distinct advantage relative to point measurement approaches such as the soil moisture probe methods. The choice of ET methods depends on project objectives and financial constraints. The eddy covariance method gives a high-resolution ET estimate for a homogeneous landscape; thus, it has the advantage for examining ecosystem processes when incorporating with simultaneous measurements of other gases (e.g., CO$_2$ and CH$_4$). However, this method is relatively costly compared with the other ET methods examined in this study. The soil water-based approach is less costly and provides a reasonable estimate of ET at a longer time scale (monthly) although it does not apply to daily ET estimates.

The early-rotation site was very sensitive to summer drought as opposed to the mid-rotation site. At the mid-rotation site, soil moisture trends demonstrated the importance of measuring water content at several depths throughout the rooting zone. Combining measurements of sapflow, soil evaporation, and canopy interception provided insights to the response of water use by plants to climatic variability and stand development. Tree transpiration and evapotranspiration were not directly comparable because understory and soil transpiration rates were not negligible. Pine transpiration showed remarkable consistency as the stands developed, regardless of the accompanying increases in LAI and changes in understory composition from early- to mid-rotation. At the early-rotation site, this conservation of water use was achieved through relatively low transpiration rates (compared with PET and ET) and compensatory transpiration among the overstory trees and understory vegetation. At this site, reduced water availability will affect trees more than understory or soil, whereas at the mid-rotation site, it would affect more understory transpiration, which might be conducive for management efforts.

**Literature Cited**


Bréda, N., R. Huc, A. Granier, and E. Dreyer. 2006. Temperate forest trees and stands under severe drought: A review of


