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Modeling Carbon and Energy Fluxes in a Heterogeneous Urban Park

February, 2015

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Kimm, Hyungsuk
Modeling Carbon and Energy Fluxes in a Heterogeneous Urban Park

어질적 도시 공원에서의 탄소와 에너지 플럭스 모델링

Under the Direction of Adviser, Prof. Ryu, Youngryel

Submitted to the Faculty of
the Graduate School of Seoul National University
December, 2014

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MEMBER
Abstract

Parks account for a large proportion of green spaces in urban regions, but most previous studies have focused on the values of recreational services in urban parks—carbon uptake by plants in urban parks has been studied less extensively. Urban parks typically form complex landscapes in space and time by integrating multiple species with open canopies. Thus, to better understand canopy photosynthesis in urban park, measuring spatial and temporal variations in photosynthetic parameters and canopy structural variables is essential. Here, we report seasonal and spatial variations in two key photosynthetic parameters ($V_{\text{cmax}}$ and $J_{\text{max}}$ which are the maximum rates of carboxylation and electron transport, respectively) and leaf area index (LAI) in Seoul Forest Park. During the peak growing season, we found an eightfold difference (20 to 149 μmol m$^{-2}$ s$^{-1}$) and fourfold difference (38 to 141 μmol m$^{-2}$ s$^{-1}$) in $V_{\text{cmax}}$ and $J_{\text{max}}$, respectively, across 10 species. Over the seasons, two woody species (Zelkova serrata and Prunus yedoensis Matsum) respectively showed three- to fivefold differences in $V_{\text{cmax}}$ and two- to fivefold differences in $J_{\text{max}}$. We evaluated whether leaf nitrogen contents could predict $V_{\text{cmax}}$ and $J_{\text{max}}$, and found significant correlations among the three variables during the peak growing season across 10 species, but no significant correlations among them over the seasons in the two woody
species. LAI computed using in-situ observations and satellite remote-sensing imagery showed a non-normal distribution with marked variation during the growing season. A sophisticated 3 dimensional model, which reflects complexity of vegetation structure in the park, well predicted carbon and energy fluxes for a day. Moreover, the model simulation with simplistic virtual scenarios clearly showed the effects of difference in tree distribution and tree size on carbon and energy fluxes (~3% and ~40%, respectively). These results highlight not only necessity of consideration of spatial and temporal variability in photosynthetic parameters and LAI for accurate estimation of canopy photosynthesis in urban parks, but also an important role of individual tree based model as a potential park design evaluating platform.

**Keywords:** leaf area index, maximum rate of carboxylation, maximum rate of electron transport, photosynthesis, 3D modeling, urban park

**Student Number:** 2013-21147
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I. INTRODUCTION

The number of urban parks is expected to increase in many urban regions due to rapid urbanization (Angel et al., 2011; Seto and Fragkias, 2005; Seto et al., 2012) and citizens’ desire for a better quality of life (Chiesura, 2004; Jim and Chen, 2006; Thompson, 2002). As urban parks account for a large proportion of green spaces in urban regions, they might partially offset carbon dioxide (CO₂) emissions in urban regions which are responsible for more than 60% of global CO₂ emissions (Birol et al., 2008; IEA, 2013). Although a few studies have attempted to quantify carbon (C) stocks in vegetation and soils in urban parks (Bae and Ryu, 2014; Hutyra et al., 2011), to our knowledge, none has quantified canopy photosynthesis, the main driver of the C cycle (Beer et al., 2010; Ryu et al., 2011).

To quantify canopy photosynthesis, an understanding of leaf-level photosynthesis is essential. The Farquhar–von Caemmerer–Berry (FvCB) model proposed a mechanistic photosynthesis paradigm (Farquhar et al., 1980), which has been widely used in predicting photosynthesis from the leaf, to global scales (dePury and Farquhar, 1997; Ryu et al., 2011; Sellers et al., 1997). The FvCB model adopts a biochemical approach based on the mechanism of the Calvin cycle, which is limited by the carboxylation rate [i.e., the amount of activated photosynthesis enzyme ribulose-1,5-
bisphosphate carboxylase/oxygenase (Rubisco)], or electron transport rate [i.e., regeneration rate of the substrate of Rubisco, ribulose bisphosphate (RuBP)]. The model computes photosynthesis separately under two assumptions: that the Calvin cycle is limited by the rate of either carboxylation or electron transport. Finally, the model provides the minimum value of the two as a photosynthesis rate. Thus, the maximum carboxylation rate \( V_{cmax} \) and the maximum electron transport rate \( J_{max} \) are the key parameters in the FvCB model and are estimated based on CO\(_2\) assimilation to leaf internal CO\(_2\) concentration curve derived from leaf gas exchange measurements (Sharkey et al., 2007). Field observations revealed that both \( V_{cmax} \) and \( J_{max} \) varied with the environment as well as season (Muraoka et al., 2010; Wang et al., 2008; Wilson et al., 2000).

Indirect estimation of the photosynthetic parameters through positive correlations between leaf nitrogen (N) contents and \( V_{cmax} \) (Amthor, 1994; Friend, 1995), and between \( V_{cmax} \) and \( J_{max} \) (Wullschleger, 1993), would require less time and effort than direct estimation from leaf gas exchange measurements. Specifically, previous studies found a positive correlation between \( V_{cmax} \) and N content per leaf area (Kattge et al., 2009). This correlation is supported by the fact that \( V_{cmax} \) is affected by the amount of the N-rich enzyme Rubisco. Also, \( V_{cmax} \) and \( J_{max} \) were found to be positively correlated. Thompson et al (1992) suggested that plants maintain a balance
between $V_{cmax}$ and $J_{max}$ by optimizing the allocation of resources (i.e., N) between Rubisco and chlorophyll to maximize photosynthesis (Wilson et al., 2000; Wullschleger, 1993). Moreover, advanced studies investigated other relationships among various leaf traits, which included leaf N concentration, leaf dry mass per leaf area (LMA), leaf N content per area, $V_{cmax}$, $J_{max}$, and the maximum rate of photosynthesis (Osnas et al., 2013; Reich et al., 1991; Wilson et al., 2000; Wright et al., 2004). These relationships, however, have not been tested in urban parks.

The leaf area index (LAI), defined as the hemi-surface leaf area per unit ground area, is the key variable to scale up photosynthesis from leaves to canopies (Baldocchi, 1997; dePury and Farquhar, 1997; Leuning et al., 1995; Ryu et al., 2011). The LAI determines mainly light interception by leaves in a canopy, which initiates canopy photosynthesis. The LAI has been measured in several forest, grassland, and cropland ecosystems (Chen, 1996; Gower and Norman, 1991; Ryu et al., 2010c). Additionally, global satellite observation systems such as the NASA Terra and Aqua satellites, have provided LAI maps over the globe at a 1-km resolution, 8-day interval (Myneni et al., 2002). However, there have been few studies that measured LAI in urban parks. Furthermore, global satellite LAI products provide 1-5-km-resolution maps, and thus the LAI in most parks, with the exception of very large parks such as Central Park in Manhattan, are unlikely to be captured by global satellite
LAI products. Thus, our understanding of spatial and temporal variations in the LAI in urban parks is limited.

Quantifying the photosynthetic parameters $V_{\text{cmax}}$ and $J_{\text{max}}$, and the LAI in urban park, is challenging due to the heterogeneity of canopy structures. To meet citizens’ diverse needs, urban parks are composed of multiple species and are spatially heterogeneous with a complex composition of land cover, such as forests, ponds, playgrounds, and lawns (Bae and Ryu, 2014; Cao et al., 2010; Jim and Chen, 2006). Heterogeneous land cover result in an uneven distribution of vegetation and makes measuring the LAI of an urban park difficult (Millward and Sabir, 2010). Moreover, a complex species composition would impose difficulties in measuring photosynthetic parameters because of species-specific differences in photosynthetic parameters (Kattge et al., 2009; Medlyn et al., 1999). Moreover, urban park vegetation is also temporally heterogeneous. The phenological patterns of factors such as the timing of leaf-out and leaf-off might vary among species and crowns (Reich et al., 1991; Wilson et al., 2000).

This study aimed to quantify the spatiotemporal heterogeneity of photosynthetic parameters and the LAI in an urban park, Seoul Forest Park, and to evaluate 3D canopy photosynthesis model against measured flux data, eventually to simulate carbon flux in various scenarios. We estimated $V_{\text{cmax}}$ and $J_{\text{max}}$ for 10 representative species during a peak growing season.
common species in the park, *Zelkova serrata* and *Prunus yedoensis Matsum*, we estimated Vcmax and Jmax across the seasons. We also measured the LAI in 10 plots over the seasons, which were combined with Landsat satellite imagery to produce spatial and temporal maps of LAI in the park. Model simulation has been conducted for a 120 m by 120 m plot by using manually measured tree profile data. Our scientific questions were as follows: 1) To what extent do photosynthetic parameters vary spatially during the peak growing season and temporally across the seasons? 2) Can Vcmax and Jmax be estimated indirectly from leaf traits data? 3) To what extent does the LAI vary across different land cover types over the seasons? 4) Can park design strategy affect carbon and energy fluxes?
II. MATERIALS AND METHODS

1. Study site

The study site is an urban park, Seoul Forest Park, in Seoul, South Korea (37.544 N, 127.038 E). The park is located beside the Han River and covers an area of 1 km². The site experiences a temperate monsoon climate. The mean annual temperature is 12.5ºC, and the mean annual precipitation is 1450 mm year⁻¹ (Korea Meteorological Administration). In 2013, the park contained over 415,000 woody plants of 95 species (Seoul Metropolitan Government).

2. Photosynthetic parameters and leaf traits measurement

To quantify the spatial and temporal variations in leaf traits, including $V_{\text{cmax}}$, $J_{\text{max}}$, LMA, and C and N concentrations, we selected 10 dominant species. Two species (Z. serrata and Pr. yedoensis) were used to estimate seasonal variations in leaf traits and eight species (Ulmus parvifolia jacq, Quercus acutissima, Quercus palustris, Betula platyphylla, Ginkgo biloba, Celtis sinensis, Pinus densiflora, and Euonymus alatus) were selected to quantify spatial variations of the leaf traits during the peak growing season (day of year, DOY 219–255). Because canopy photosynthesis models require sunlit leaf traits as input parameters, we sampled only sunlit leaves located at the...
top of crowns or the outer side of the south face of isolated crowns (dePury and Farquhar, 1997; Kattge et al., 2009; Ryu et al., 2011). To sample sunlit leaves, we used 3-m-long pruning scissors. We chose one tree per one species and collected three branches each of which held at least three leaves. One leaf of each branch was used for leaf gas exchange measurements and all three leaves of the branch were transported to the laboratory for measurement of LMA, and leaf N and C concentrations.

We estimated the key photosynthetic parameters, \( V_{c_{\text{max}}} \) and \( J_{\text{max}} \), from leaf gas exchange measurements using a portable photosynthesis system (Li-6400; Li-Cor Inc., Lincoln, NE, USA). First, we measured the photosynthesis rate (A) responding to leaf internal CO\(_2\) concentrations (Ci) to obtain A/Ci curves. For each leaf, we used the automated program mode in Li-6400 to create a A/Ci curve by changing leaf external CO\(_2\) concentrations (400, 200, 50, 100, 200 … 1400 ppm). Second, we estimated \( V_{c_{\text{max}}} \) and \( J_{\text{max}} \) from the obtained A/Ci curves using a least-squares curve-fitting method to fit the measured A/Ci curve to the FvCB model. We estimated \( V_{c_{\text{max}}} \) and \( J_{\text{max}} \) using the Microsoft Excel\textsuperscript{TM} spreadsheet provided by Sharkey et al. (2007). We report \( V_{c_{\text{max}}} \) and \( J_{\text{max}} \) values that were corrected to the leaf temperature at 25ºC.

To quantify LMA and foliar C and N concentrations, we scanned all sampled leaves and computed leaf areas using the MATLAB image-processing toolbox (MathWorks Inc., Woburn, MA, USA). Then we
measured the dry mass of each leaf using a high-precision scale (accuracy: 0.001 g, CUX220H; CAS Corp., Seoul, South Korea) after 48 h of oven-dry at 80.0°C (C-DH; Chang Shin Scientific Co., Pusan, South Korea). Dried leaves were ground into powder to estimate C and N concentrations using an Elemental Carbon Analyzer (Flash EA 1112; Thermo Electron Co., Waltham, MA, USA) at the National Instrumentation Center for Environmental Management (NICEM) of Seoul National University.

3. Leaf area index measurement

3.1. Plot design

We randomly established 10 20 × 20-m plots in the planted areas in the park. Ten plots represented one evergreen needleleaf forest (ENF), one mixed forest (MF), one deciduous needleleaf forest (DNF), and seven deciduous broadleaf forests (DBFs). The plots showed a tenfold difference in stem density and twofold difference in tree height. The stem density, mean tree height, and species composition of each plot are shown in Table 1.
<table>
<thead>
<tr>
<th>Plots</th>
<th>STEM DENSITY (stem/m²)</th>
<th>TREE HEIGHT (m)</th>
<th>SPECIES</th>
</tr>
</thead>
<tbody>
<tr>
<td>DBF1</td>
<td>0.09</td>
<td>8</td>
<td><em>Acer palmatum</em> Thumb, <em>Quercus mongolica</em></td>
</tr>
<tr>
<td>DBF2</td>
<td>0.13</td>
<td>9</td>
<td><em>Quercus acutissima</em>, <em>Acer Buegerianum</em></td>
</tr>
<tr>
<td>DBF3</td>
<td>0.07</td>
<td>15</td>
<td><em>Platanus acerifolia</em></td>
</tr>
<tr>
<td>DBF4</td>
<td>0.12</td>
<td>11</td>
<td><em>Quercus acutissima</em>, <em>Ginkgo biloba</em>, <em>Acer buegerianum</em>, <em>Aesculus turbinata</em></td>
</tr>
<tr>
<td>DBF5</td>
<td>0.12</td>
<td>8</td>
<td><em>Quercus acutissima</em>, <em>Prunus yedoensis Matsum</em>, <em>Zelkova serrata</em>, <em>Ulmus parvifolia jacq</em></td>
</tr>
<tr>
<td>DBF6</td>
<td>0.04</td>
<td>8</td>
<td><em>Zelkova serrata</em>, <em>Carpinus laxiflora</em>, <em>Acer palmatum</em></td>
</tr>
<tr>
<td>DBF7</td>
<td>0.04</td>
<td>9</td>
<td><em>Zelkova serrata</em></td>
</tr>
<tr>
<td>DNF</td>
<td>0.39</td>
<td>11</td>
<td><em>Ginkgo biloba</em></td>
</tr>
<tr>
<td>MF</td>
<td>0.05</td>
<td>11</td>
<td><em>Pinus strobus</em>, <em>Ulmus parvifolia jacq</em>, <em>Metasequoia glyptostroboideis</em>, <em>Liriodendron tulipifera</em></td>
</tr>
<tr>
<td>ENF</td>
<td>0.08</td>
<td>14</td>
<td><em>Pinus densiflora</em></td>
</tr>
</tbody>
</table>
3.2. Digital cover photography (DCP)

We used DCP to estimate the LAI (Macfarlane et al., 2007; Ryu et al., 2014; Ryu et al., 2012). Photographs were obtained at 1-m height under canopies using a digital single-lens reflex camera (Nikon 600D; Nikon, Tokyo, Japan). We obtained three to seven photographs per plot viewing toward 57º from the zenith direction. G-function is the projection coefficient of unit foliage area on a plane perpendicular to the view direction and at a 57º-view zenith angle; the G-function is 0.5 regardless of leaf inclination angle distributions (Nilson, 1971; Wilson, 1960). We chose the 57º-view zenith angle because measuring leaf inclination angles from the canopy top to bottom for various species was difficult. The lens had focal length of 28.8 mm (35-mm equivalent). We obtained photographs under the aperture priority mode and low aperture value (>f/1/5.6) to broaden the depth of field within photographs (Pentland, 1987). We also set the shutter speed to shorter than 1/15 s to avoid blurred photographs. We chose the proper exposure level manually by checking the histogram of the blue channel using a built-in camera liquid-crystal display (LCD) to minimize overexposed canopy pixels for accurate estimation of the LAI. Each time we visited the field, we obtained photographs at the same positions and in the same directions to minimize uncertainties caused by inconsistent sampling footprints. We obtained photographs every 2 weeks on average from DOY 121 to DOY 333.
To estimate the LAI from photographs, we first extracted the blue channel, which shows the most marked contrast between the sky and vegetation. In the blue channel histogram, we separated pixels into sky pixels or vegetation pixels using a two-corner method to set the thresholds for pixel classification (Macfarlane, 2011). The obtained binary images enabled us to calculate the fraction of crown cover (CC) and gap fraction within crowns; i.e., crown porosity (CP; Macfarlane et al., 2007). Thus, we finally calculated the LAI using the modified Beer’s law (Ryu et al., 2012):

$$\text{LAI} = \frac{-CC \times \log(CP)}{k} \gamma,$$  

(2)

where $k$ is the light extinction coefficient [$G(57°)/\cos(57°)$] and $\gamma$ is the needle-to-shoot area ratio. We computed the LAI for individual photographs then averaged the LAI values in one plot to account for foliar clumping effects (Ryu et al., 2010b). In the case of needleleaf species ($P. densiflora$ and $Pinus strobus$), we measured the needle-to-shoot area ratio to account for foliar clumping effects appearing at the shoot scale as proposed by Ryu et al. (2014). The $\gamma$ value for $P. densiflora$ was $2.11 \pm 0.28$. The image analysis was conducted with a technical computing language, MATLAB (MathWorks Inc.).
4. Satellite remote-sensing data processing

We used Landsat 8 imagery (30-m resolution). We selected satellite data for 5 days (DOY 86, 131, 179, 259, and 355) that had less than 10% cloud cover per image. To remove path radiance effects in the atmosphere, we adopted the dark object subtraction method (Song et al., 2001). To obtain corrected digital numbers after the atmospheric correction, we converted the digital numbers to spectral reflectance using conversion coefficients provided by the Landsat 8 metafile. We computed the normalized difference vegetation index (NDVI) using red and near-infrared reflectance (Tucker, 1979):

\[
NDVI = \frac{\rho_{NIR} - \rho_{RED}}{\rho_{NIR} + \rho_{RED}},
\]

where \(\rho_{NIR}\) and \(\rho_{RED}\) are the spectral reflectance of near-infrared and red, respectively. We extracted NDVI values within each LAI measurement plot. To infer the LAI from the NDVI (Ryu et al., 2010a), we developed an exponential regression model (LAI = 3.44 \(\exp(NDVI)\) – 3.38; \(R^2 = 0.79, P < 0.001\]). Using this equation, we generated LAI maps from Landsat NDVI imagery for 5 days (DOY 86, 131, 179, 259, and 355).
5. Model simulation

We used FLiES combined with CANOAK to simulate carbon and energy fluxes based on 3D radiative transfer simulation. We collected tree position and profile data to construct individual trees in model simulation. Only DOY 153 has been chosen because it was the clearest day (i.e., clear sine curve in incoming shortwave radiation) with stable wind condition (the smallest footprint of the flux measurements) during the summer season in 2013.

5.1. Model description

A 3D radiative transfer model, FLiES simulated radiative transfer by a Monte Carlo ray tracing which is randomly simulating the path length of each of the 22.5 million photons based on probability of a scattering event. Simulated amounts of absorbed photons at three different range of wavelength, photosynthetically active radiation, near infrared, thermal infrared, were used to simulate photosynthesis, evapotranspiration, and skin temperature at different surfaces.

5.2. Collecting crown data

We measured tree size and position data through a series of field work. Positions of trees, tree height, crown depth, crown width, and diameter at breast height (DBH) have been measured within a 120 m by 120 m size plot.
We used typical tapes to measure DBH and crown width. For crown width measurement, we measured distance from a trunk to the end of the crown at four directions (0, 90, 180, 270), and then averaged 4 values. To obtain vertical size of trees, we used a laser meter (Leica DISTO™ D5, Leica Geosystems, Hexagon, Sweden) at distant location from target trees to measure distance and angles toward at specific points of trees. Through a simple calculation, we estimated tree height and crown depth. Tree positions were captured by using distances from two reference points.

5.3. Flux measurement

We measured carbon and water fluxes, and meteorological variables (air temperature, humidity, wind speed and direction, incoming shortwave radiation) with the Eddy covariance measurements on the top of a building in the park. We obtained 30 min averaged data of incoming shortwave radiation, wind speed and direction, carbon and water fluxes, air temperature, and humidity.

5.4. Simulations with virtual scenarios

We developed two sets of scenarios with difference in tree size and tree distribution. With the same set of trees, three scenarios, SCN_1, SCN_2, SCN_3, are composed with different tree distributions. In case of scenarios
with different tree sizes (SCN_L, SCN_M, SCN_MX, SCN_S), we adopted regular distributions of trees to only test the effect of different tree sizes with the same leaf area density and leaf area. Individual tree elements and the compositions are presented in Table 2.

Table 2. Description of scenarios for model simulation. LAD, LAI, CC represents leaf area density (m$^2$/m$^3$), leaf area index (m$^2$/m$^2$), crown cover (%), respectively. “ELEMENTS” shows the profile of individual trees in scenarios.

<table>
<thead>
<tr>
<th>NAME</th>
<th>SCN_1</th>
<th>SCN_2</th>
<th>SCN_3</th>
<th>ELEMENTS</th>
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<td>IMAGE</td>
<td>![Image]</td>
<td>![Image]</td>
<td>![Image]</td>
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</tr>
<tr>
<td>LAD</td>
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<td>1.623</td>
<td>1.623</td>
<td></td>
</tr>
<tr>
<td>LAI</td>
<td>1.51</td>
<td>1.49</td>
<td>1.46</td>
<td></td>
</tr>
<tr>
<td>CC</td>
<td>55.77</td>
<td>47.20</td>
<td>54.15</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>NAME</th>
<th>SCN_L</th>
<th>SCN_M</th>
<th>SCN_MX</th>
<th>SCN_S</th>
</tr>
</thead>
<tbody>
<tr>
<td>IMAGE</td>
<td>![Image]</td>
<td>![Image]</td>
<td>![Image]</td>
<td>![Image]</td>
</tr>
<tr>
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<td>1.623</td>
<td>1.623</td>
<td>1.623</td>
<td>1.623</td>
</tr>
<tr>
<td>LAI</td>
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<td>2.92</td>
<td>2.91</td>
<td>2.99</td>
</tr>
<tr>
<td>CC</td>
<td>21.39</td>
<td>32.19</td>
<td>37.50</td>
<td>53.43</td>
</tr>
</tbody>
</table>
III. RESULTS AND DISCUSSION

1. To what extent do photosynthetic parameters vary spatially during the peak growing season and temporally across the seasons?

Two photosynthetic parameters, $V_{\text{cmax}}$ and $J_{\text{max}}$, of the 10 species in the park presented a wide range of variation during the peak growing season [7.5-fold variation (from 20 to 149 μmol m$^{-2}$ s$^{-1}$) in $V_{\text{cmax}}$ and 3.7-fold variation (from 38 to 141 μmol m$^{-2}$ s$^{-1}$) in $J_{\text{max}}$; Fig. 1]. *Ulmus parvifolia jacq* and *C. sinensis* showed the highest and lowest $V_{\text{cmax}}$ values (149 and 20 μmol m$^{-2}$ s$^{-1}$, respectively), *E. alatus* and *C. sinensis* displayed the highest and lowest $J_{\text{max}}$ values (141 and 38 μmol m$^{-2}$ s$^{-1}$, respectively), and *U. davidiana* and *C. sinensis* had the highest and lowest values in the photosynthetic parameters. However, we observed that the species pairs were co-located within a 5-m distance in several places. In a meta-analysis study, Kattge et al. (2009) reported 57.7 ± 21.2 μmol m$^{-2}$ s$^{-1}$ [mean ± 1 standard deviation (S.D.), $n = 404$] $V_{\text{cmax}}$ in temperate broad-leaved deciduous trees globally, while our results indicated 82.9 ± 37.3 μmol m$^{-2}$ s$^{-1}$ in $V_{\text{cmax}}$ ($n = 7$) for the same plant functional type trees, indicating greater variation in $V_{\text{cmax}}$.

The photosynthetic parameters of *Z. serrata* and *Pr. yedoensis* showed large seasonal variations (Fig. 2). Over the seasons, $V_{\text{cmax}}$ varied from 20 to 100
μmol m$^{-2}$ s$^{-1}$ and 40 to 120 μmol m$^{-2}$ s$^{-1}$ for Z. serrata and Pr. yedoensis,

Fig. 1. (a) $V_{cmax}$ and (b) $J_{max}$ in 10 species measured during the peak growing season. Error bars indicates 95% CI.
respectively. The trends depicted leaf ontogeny and senescence, as reported for deciduous broad-leaved trees in previous studies (Muraoka et al., 2010; Wang et al., 2008; Wilson et al., 2000). Our results showed prolonged peak $V_{c_{\text{max}}}$ values until DOY 250, followed by a reduction, which is similar to the findings of a study conducted in a temperate deciduous forest in Japan (Muraoka et al 2010).

The heterogeneous composition of plant species in urban parks, which caused marked variations in photosynthetic parameters, requires a different approach to canopy photosynthesis modeling. In natural ecosystems, high biodiversity of woody plants under an open canopy, which is the case in urban parks, is uncommon. For example, savanna is a typical case of open canopy but includes lower woody plant biodiversity, which are distributed sparsely (Baldocchi et al., 2004). In contrast, tropical forests have high woody plant biodiversity, which represent a closed canopy (Asner et al 2009). The large variation in $V_{c_{\text{max}}}$ during the peak growing season in the urban park (20–149 μmol m$^{-2}$ s$^{-1}$, nine species of woody plants) was greater than that reported previously in an Amazon tropical forest (30–80 μmol m$^{-2}$ s$^{-1}$, 12 species; Domingues et al., 2012), a temperate deciduous forest (40–60 μmol m$^{-2}$ s$^{-1}$, 4 species; Wilson et al 2000), and an woody savanna forest (130 μmol m$^{-2}$ s$^{-1}$, 1 species; Baldocchi et al., 2004). Therefore, we assume that an individual-tree-based three-dimensional (3D) canopy photosynthesis model is more appropriate for estimating canopy photosynthesis in urban parks.
Conventional photosynthesis models—such as a big-leaf model (Sellers 1985) or two-leaf models (dePury and Farquhar 1997, Ryu et al 2011)—which simplify ecosystem structure and functions into one or two big leaves, are unlikely to simulate accurately canopy photosynthesis in urban parks.

Fig. 2. Seasonal variation in (a) V_{cmax} and (b) J_{max} for Zelkova serrata and Prunus yedoensis Matsum. Error bars indicate 95% CI.
2. Can $V_{c\text{max}}$ and $J_{\text{max}}$ be estimated indirectly from leaf traits data?

We confirmed the possibility of estimating the photosynthetic parameters from leaf N content per unit leaf area for data measured during the summer (DOY 207–255) across the species. These data showed significant correlations between $V_{c\text{max}}$ and $J_{\text{max}}$ ($R = 0.85, P < 0.001$; Fig. 4), and between $V_{c\text{max}}$ and N content ($R = 0.7, P < 0.001$; Fig. 3). The slope of the linear relationship between $V_{c\text{max}}$ and $J_{\text{max}}$ (0.76) was lower than those reported previously; i.e., 1.2–3.33 (Grassi et al., 2005; Wilson et al., 2000; Wullschleger, 1993).

![Graph showing comparison of leaf nitrogen content per unit area and $V_{c\text{max}}$ for 10 species measured during the peak growing season.](image)

**Fig. 3.** Comparison of the leaf nitrogen content per unit area and $V_{c\text{max}}$ for 10 species measured during the peak growing season.
Fig. 4. Comparison of $V_{c\text{max}}$ and $J_{\text{max}}$ for 10 species measured during the peak growing season.

However, we found no significant correlation between $V_{c\text{max}}$ and N content for the data on $Z. \text{serrata}$ and $Pr. \text{yedoensis}$ measured over the seasons ($R = 0.07, P = 0.16$). We speculate that the summer monsoon decouples the $V_{c\text{max}}$ and N relationship. The study year (2013) experienced a remarkably long monsoon period (DOY 168–216). We assume that both the low-light environment caused by frequent clouds and rainfall would decrease the leaf N content during the monsoon period (34 days of precipitation with 20 mm
day\(^{-1}\) on average according to a weather station located 1 km from the study site; Korea Meteorological Administration). Continuous rainfall might have reduced leaf N contents because N leaching could account for \(\sim 15\%\) of the total amount of N returned from the leaves to soils (Chapin and Moilanen, 1991; Lambers et al., 1998). Indeed, our results showed that leaf N contents in the two species decreased significantly during the monsoon period (shaded area in Fig. 5), whereas the \(V_{cmax}\) values of both species were fairly stable during that period (Fig. 2). Furthermore, during autumn, leaf N contents increased (Fig. 5), which is contrary to the nutrient resorption pattern (i.e., reduction of leaf N contents due to leaf nutrient withdrawal by plants before leaf abscission; Reich et al. 1991; Wilson et al. 2000). We do not have strong evidence for the uncommon pattern of leaf N seasonality at this study site. We speculate that a prolonged monsoon period and sufficient N input to the urban park through atmospheric N deposition and fertilization might be related to the leaf N seasonality.

We found a significant correlation between \(V_{cmax}\) and \(J_{max}\) for \(Z.\) serrata and \(Pr.\) yedoensis across the seasons \((R = 0.85, P < 0.01)\). Thus, the tight correlations between \(V_{cmax}\) and \(J_{max}\) were valid both spatially during the peak growing season and temporally across seasons. Thus, finding correlations between leaf N contents and \(V_{cmax}\) is the essential step. Leaf N contents were predictive of \(V_{cmax}\) across 10 species during the peak growing season, but not
for the two species across the seasons. Identification of correlations between $V_{cmax}$ and other variables—such as the LAI (Fig. 6)—over the seasons might offer an alternative method of inferring seasonality in $V_{cmax}$ (Houborg et al., 2009; Ryu et al., 2011), although we were unable to test this hypothesis because the crowns from which the Z. serrata and Pr. yedoensis leaf samples were collected were not included in our LAI observation plots.

![Seasonal trend in leaf nitrogen content per unit area for Zelkova serrata and Prunus yedoensis Matsum. Error bars indicate 95% CI. The shaded area represents the monsoon period.](image)

Fig. 5. Seasonal trend in leaf nitrogen content per unit area for *Zelkova serrata* and *Prunus yedoensis* Matsum. Error bars indicate 95% CI. The shaded area represents the monsoon period.
3. To what extent does the LAI vary across different land cover types over the seasons?

The peak LAI values derived from DCP varied from 3.1 to 4.4 across the plots in the park during the peak growing season (Fig. 6). All plots showed a clear seasonality in the LAI. We obtained the spatial distribution of the LAI for the entire park by merging in situ LAI observations using DCP and Landsat NDVI images (Fig. 7a). The LAI clearly showed considerable spatial variation from 1.03 to 3.76 at DOY 259.

![Leaf area index variation across different land cover types](image)

Fig. 6. Seasonal variation in the leaf area index in 10 plots.
We investigated the histograms of the LAI in the park for DOY 86, 131, 179, 259, and 355 (Fig. 7b). During winter, the histogram of the LAI showed the lowest mean ± S.D. values (DOY 86: 0.97 ± 0.15, DOY 355: 0.60 ± 0.13). The skewness and kurtosis were close to 0 and 3, respectively, indicating that the histograms followed a normal distribution. In the peak growing season (e.g., DOY 259), the LAI values showed the largest spread in distribution (0.55 S.D.) and a non-normal distribution caused by higher kurtosis (4.49), which was skewed left (−1.20). The non-normal, considerable spread in the LAI distribution reflects the heterogeneous canopy structure in the urban park, which cannot be represented by a single mean value.

Given the large spatial and temporal variations in photosynthetic parameters and LAI, we argue that individual tree-based 3D canopy photosynthesis models such as FLiES (Kobayashi and Iwabuchi, 2008) and MAESTRA (Wang and Jarvis, 1990) will be essential for predicting canopy photosynthesis in urban parks. We acknowledge that the 30-m resolution in Landsat images might be insufficient to capture spatial heterogeneity in urban parks. Recent advances in light detection and ranging (LiDAR) offers new opportunities to extract 3D canopy structure information, which will facilitate quantification of the complex canopy structures in urban parks.
Fig. 7. (a) Leaf area index (LAI) map for day of year (DOY) 259 derived from Landsat images. (b) Histograms of Landsat-derived LAI values in the park for DOY 86, 131, 179, 259, and 355. The curves indicate histograms fit to a normal distribution.
4. Can park design strategy affect carbon and energy fluxes?

The model simulated carbon and energy fluxes were accurate compared to the eddy covariance measurement results (R squared values were 0.77, 0.76, 0.61 for latent- and sensible heat fluxes, and net ecosystem exchange, respectively; Fig. 9). Considering that the nighttime footprint of the flux measurement would expand beyond the canopies, thus to include roads with heavy traffic, mismatch between the simulation result and the measurement could be partially explained with carbon emission from the traffic. Negative anomalies in model simulated latent heat flux might be resulted from none of consideration of water surface although the simulated plot includes a small pond. This, in turn, caused positive anomalies in sensible heat flux with energy balance algorithm (Kyaw Tha Paw, 1987).

Fig. 8. Satellite image (left), and visualized input data (right) of the model simulation plot site (120 m by 120 m). Circles in both plots represents the footprint of the eddy covariance measurement.
Fig. 9. Comparison between the simulated- and measured carbon and energy fluxes in the plot. Extreme values with numerical errors are manually eliminated.

Under simplistic scenarios, the model simulated results showed that carbon and energy fluxes change with difference in park vegetation composition (Fig. 10, Fig. 11). Difference in tree distribution lead to little difference (less than 10%) in both carbon and energy fluxes (Fig. 10), but difference in tree size significantly affected the results (~ 40%). We suspect the effect of the difference in crown cover of the scenarios with different tree sizes would have been dominant as higher crown cover largely affect absorbed radiation, which could lead to large difference in photosynthesis and energy partitioning. Contrast results from the two sets of simulations clearly show what factor
is effective and what is not, even in this simplistic simulations.

Fig. 10. Difference of results between SCN_1 and SCN_2, and SCN_1 and SCN_3. See Table 2 for scenarios in the legend.

Fig. 11. Difference of results between SCN_L and SCN_M, SCN_L and SCN_MX, and SCN_L and SCN_S. See Table 2 for scenarios in the legend.
5. **Broader implications for future urban park design**

Urban planners and designers have paid less attention to the potential role in carbon uptake by plants. Mixture of open space and clumped canopy with diverse species, which promotes diverse activities by visitors, pervades in many urban parks. As shown in the results, the study site included a range of different species with large spatial and temporal variability in canopy structure and photosynthetic parameters (Figure 1 and 7). Canopy photosynthesis generally increases with light when solar irradiance is low, and it becomes saturated when solar irradiance is high. Trees that have high Vcmax (i.e. a proxy of photosynthetic capacity) could increase canopy photosynthesis even in high solar radiation conditions. Thus, archiving Vcmax values in typical urban park tree species will be useful in selecting tree species in planting design. For example, in isolated, sparsely distributed, or row-planted canopies, planting trees with higher Vcmax which could more efficiently use high level of solar irradiance would be desirable to enhance canopy photosynthesis. We do not argue carbon capture is the most important function in urban parks; rather, we hope carbon sequestration could be harmonized in the ecosystem services that urban parks provide. Owing to 3D canopy photosynthesis models, it is possible to simulate carbon fluxes with a range of different scenarios that include different plant distributions, plant species, canopy heights and multi-layered canopy structures. We expect
further applications of 3D canopy photosynthesis model into urban parks will be warranted in the future studies to better understand carbon cycles in urban regions as well as to help simulation-informed park planning and design.
IV. Summary and conclusions

In this report, we described the spatial and temporal patterns in photosynthetic parameters ($V_{c\text{max}}$ and $J_{\text{max}}$) and the LAI in an urban park in Seoul. The answers to the three scientific questions are as follows: (1) To what extent do photosynthetic parameters vary spatially during the peak growing season and temporally across the seasons? During the peak growing season, we found an eightfold difference in $V_{c\text{max}}$ and fourfold difference in $J_{\text{max}}$ across 10 species. Over the seasons, two woody species ($Z. \text{serrata}$ and $P. \text{yedoensis}$) showed three- to fivefold differences in $V_{c\text{max}}$ and two- to fivefold differences in $J_{\text{max}}$, respectively. (2) Can one estimate $V_{c\text{max}}$ and $J_{\text{max}}$ indirectly from leaf traits data? We found that the leaf N content was predictive of $V_{c\text{max}}$ across 10 species during the peak growing season. Also, a strong positive correlation existed between $V_{c\text{max}}$ and $J_{\text{max}}$. However, across the seasons, the leaf N content was not significantly correlated with $V_{c\text{max}}$, probably because of the prolonged summer monsoon. Thus, we did not find a universal relationship among leaf N, $V_{c\text{max}}$, and $J_{\text{max}}$. (3) To what extent does the LAI vary across land cover types over the seasons? LAI maps derived from the Landsat NDVI and in situ LAI observations revealed a normal distribution with a small spread in LAI values in winter, whereas a non-normal distribution with a large
spread in LAI estimates was observed during the peak growing season. (4) Can park design strategy affect carbon and energy fluxes? Model validation against the measurement proved reliable model performance, and further simulations revealed significant differences from different tree size, but not from different tree distribution implicating the model simulation could be used for evaluating park designs in terms of carbon and energy fluxes. Our findings highlight the possibilities of large spatial and temporal variation in photosynthesis, and the potential effects of park designs on carbon and energy fluxes in urban parks. Thus, we conclude that the use of an individual-tree-based, 3D canopy photosynthesis model is essential for accurately predicting canopy photosynthesis, and is capable of evaluating park designs in terms of carbon budget and energy partitioning.
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국문 초록

도시 공원은 도시 지역 내에서의 상당한 비율을 차지하지만, 기존의 도시 관련 연구들은 도시 공원 식생의 탄소 흡수에 대해서는 연구가 많이 이뤄지지 않았다. 일반적인 도시 공원은 다양한 수종을 포괄하고, 개방적 수관부의 배치를 통해 시간적, 공간적으로 매우 복합적인 경관을 구성한다. 따라서, 도시 공원에서의 광합성을 보다 잘 이해하기 위해서는 광합성 매개변수와 수관 구조 변수의 공간적, 시간적 차이를 관측하는 것이 필수적이다. 본 연구에서는 서울 숲 공원을 대상으로, 두 개의 핵심적인 광합성 매개변수 (최대 카르복시화율과 최대 전자 전달률)를 나타내는 $V_{\text{cmax}}$ 와 $J_{\text{max}}$ 와 엽면적지수의 계절적, 공간적 차이를 보인다. 대표 성장 시기에 10개의 수종 간에 $V_{\text{cmax}}$ 와 $J_{\text{max}}$ 는 각각 8배, 4배의 차이를 보였다. 계절적으로는 두 대표 수종인 느티나무와 벚나무가 $V_{\text{cmax}}$ 에서는 각각 3배, 5배의 차이를, $J_{\text{max}}$ 에서는 2배, 5배의 차이를 보였다. 본 연구는 또한 임의 질소 함량을 통해 두 매개변수 추정이 가능한 지를 평가했고, 최대 성장기의 10개 수종 대상의 관측자료에서는 질소 함량과 두 매개 변수 사이에서의 상당한 상관관계를 확인했으나, 두 수종의 여러 계절 간 관측자료를 분석한 결과에서는 상관관계를 특정 지을 수 없었다. 엽면적 지수는 현장 관측과 위성 원격 탐사 영상을 통해 계산했으며, 시간적, 공간적으로 비정규분포를 보이며 큰 차이를 나타냈다. 정교하게 도시 공원 내 복합적인 구조적 특성을 반영한 3차원 모델은 탄소와 에너지 플럭스를 비교적 정확히 추정했다. 또한, 가장 시나리오를 활용한 모델 시뮬레이션은 수목의 분포와 개별 수관의 크기에 따른 효과를 명확히
보여주었다. 수목 분포 차이는 탄소 및 에너지 플럭스 결과에서 최대 3%의 차이만을 보였고, 반면에 개별 수관의 크기 차이는 최대 40%의 큰 차이를 보였다. 본 결과는 각각의 요소가 탄소수지 및 에너지 분할에 미칠 수 있는 효과를 평가할 수 있는 도구임을 시사한다. 위에서 제시된 본 논문의 결과들은 정확한 광합성 추정을 위해 광합성 매개변수 및 엽면적 지수의 공간적, 시간적 변이를 고려함이 중요함을 부각시킬 뿐 아니라, 공원 설계인 평가 플랫폼으로서의 3차원 모델링의 잠재적 역할을 시사한다.

주요어: 엽면적지수, 최대 카르복시화율, 최대 전자 전달률, 광합성, 3차원 모델링, 도시공원

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