Reducing the Excessive Evaporative Demand Improved the Water-use Efficiency of Greenhouse Cucumber by Regulating the Trade-off between Irrigation Demand and Plant Productivity

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Abstract. Although atmospheric evaporative demand mediates water flow and constrains water-use efficiency (WUE) to a large extent, the potential to reduce irrigation demand and improve water productivity by regulating the atmospheric water driving force is highly uncertain. To bridge this gap, water transport in combination with plant productivity was examined in cucumber (Cucumis sativus L.) grown at contrasting evaporative demand gradients. Reducing the excessive vapor pressure deficit (VPD) decreased the water flow rate, which reduced irrigation consumption significantly by 16.4%. Reducing excessive evaporative demand moderated plant water stress, as leaf dehydration, hydraulic limitation, and excessive negative water potential were prevented by maintaining water balance in the low-VPD treatment. The moderation of plant water stress by reducing evaporative demand sustained stomatal function for photosynthesis and plant growth, which increased substantially fruit yield and shoot biomass by 20.1% and 18.4%, respectively. From a physiological perspective, a reduction in irrigation demand and an improvement in plant productivity were achieved concomitantly by reducing the excessive VPD. Consequently, WUE based on the criteria of plant biomass and fruit yield was increased significantly by 43.1% and 40.5%, respectively.

Water shortage is the bottleneck for sustainable agricultural production, particularly in semiarid regions. An abrupt decline in the supply of irrigation water to farmers has aroused growing concern, and irrigation management has shifted from emphasizing the production per unit area toward maximizing the production per unit of water consumed (Deng et al., 2006). From classic views of physics, water transport along the soil–plant–atmosphere continuum is a passive process driven by gradients of free energy. The driving force for water movement is determined by the water potential gradient in the liquid phase (from soil to leaf) and the difference in VPD along the gas phase (from the internal leaf to the atmosphere) (Fricke, 2016; Wheeler and Stroock, 2008).

Regulation of the soil water status has long been recognized as a useful way to improve WUE. Water-saving irrigation techniques such as controlled, alternate partial root-zone irrigation and deficit irrigation have been widely applied for crop production as a result of the reduced water supply without a significant yield reduction (Gheysari et al., 2017; Kang and Zhang, 2004; Yang et al., 2017). Although atmospheric VPD mediates water flow and constrains water productivity to a large extent, regulation of the atmospheric water driving force receives far less attention from growers. The potential to reduce plant irrigation consumption and improve WUE by regulating the atmospheric driving force is highly uncertain.

The increasing sophistication of controlled-environment agriculture facilitates atmospheric environmental regulation (Katsoulas et al., 2001). Because VPD is an indicator of the atmospheric driving force for water movement, greenhouse operations are moving toward controlling evaporative demand according to VPD (Lu et al., 2015; Zhang et al., 2015). Previous studies have suggested that greenhouse VPD regulation can maintain optimal ranges of temperature and relative humidity (RH) simultaneously, and therefore effectively enhance plant photosynthesis and growth (Lu et al., 2015; Zhang et al., 2015, 2017, 2018). It is generally considered that the movement of water through the soil–plant–atmosphere continuum is driven by a gradient in water potential in the liquid phase (from soil to leaf apoplast) and a difference in the water vapor pressure in the gas phase (from leaf to atmosphere). Therefore, VPD control was hypothesized to perform significant roles in reducing irrigation demand and moderating plant water stress, which would enhance plant photosynthesis and improve WUE. Cucumber is one of the major vegetable crops cultivated under greenhouse conditions and is considered one of the most important economic vegetables. Cucumber plants require large amounts of water, especially during the flowering and fruiting periods, as the water content of fruit exceeds 90%. Despite the existence of information about the effect of soil moisture on irrigation water demand, few studies have focused on examining the relationship between VPD and irrigation water demand in greenhouse cucumber production. The current study provided innovative practices in saving water consumption and improving WUE in greenhouse cucumber production. The processes of carbon gain and water consumption were coupled to fill the gap between VPD regulation and WUE in greenhouse cucumber production.

Materials and Methods

Plant materials and growth conditions. Experiments were conducted in two outdoor chambers with similar characteristics (length, 10 m; width, 5 m; height, 3.5 m) during the spring–summer seasons from April to July. The variability in environmental factors...
between the two chambers was examined before the experiments. No significant differences in environmental factors were observed between the two chambers. The roof ventilation systems in the two treatments were activated when the temperature inside exceeded the optimum range for cucumber growth.

Cucumber (cv. Jinyou) plants were transplanted at the four-leaf stage and grown in white pots (diameter, 40 cm; height, 30 cm). One plant was grown in each pot, and each pot was filled with the same amount of garden mix substrate containing an organic substrate-perlite mixture (3:1). The planting density was four plants per m. All axillary shoots were removed, and fruit pruning was carried out to leave only a single fruit per node.

The soil surface was covered with a circular polythene sheet to prevent soil water evaporation. Plant transpiration was measured by a standardized gravimetric approach of daily pot weighing with an electronic balance, as described in previous research (Kadam et al., 2015). Excluding soil evaporation, the difference between pot weighing can be considered the plant-transpired water consumption. Soil moisture was maintained uniformly at 85% to 90% of field capacity by adding the exact amount of water needed to bring back the moisture content to the desired target.

**Experimental design.** Two environmental chambers were controlled to have the same growth conditions but a contrasting VPD. A high VPD was maintained in natural conditions without environmental regulation; a low VPD was achieved by artificial humidification when evaporative demand exceeded the optimal range. In the low-VPD compartment, humidification was controlled using a fogging system (spray pressure, 2–6 MPa; droplet size, 25.8–46.2 μm) with a binary fluid mist nozzle. Spraying was activated automatically when the greenhouse VPD exceeded 1.5 kPa, which is the recommended value for horticultural crops (Bakker, 1990; Iraqi et al., 1995). The fogging system operated continuously and was turned off when VPD decreased to less than the set point. A randomized complete block design was adopted.

**Environmental measurements.** Air temperature, relative humidity (RH), and light intensity were monitored by sensors (ZDR-20; WuGe Instruments Co., Ltd., China) installed ≈2.5 m above the ground in the center of each greenhouse. VPD was calculated from Ta and RH.

**Measurements of leaf gas exchange.** Leaf gas exchange parameters were measured with a portable photosynthesis system (LI-6400; LI-COR Inc., Lincoln, NE) ≈40 d after transplanting (from 0900 to 1200 HR). All samples were the youngest and fully expanded leaves at the same nodes of the plants. Three leaves per plant were sampled for the measurements. The environmental conditions in the leaf chamber were set close to the open-field conditions in the greenhouse.

The diurnal changes in plant transpiration rate were determined by weighing the pots hourly. The plant transpiration rate was calculated as the ratio of the pot weight change vs. the time between two weighings.

Determination of the net photosynthetic rate and stomatal conductance (gs) was repeated with 10 plants in each treatment, and a total of three measurements per plant were performed after steady state and equilibration were attained. Two photosynthesis systems were used to enable simultaneous gas exchange measurements. Stomatal limitation (Ls) was estimated according to $L_s = 1 - C_i/C_a$, where $C_i$ is the intercellular carbon dioxide (CO2) concentration and $C_a$ is the ambient CO2 concentration (Duan et al., 2014). Instantaneous WUE (InstWUE) at the leaf scale was calculated as the amount of carbon gain in photosynthesis rate ($P_n$) per unit of water transpired ($T_i$), $\text{InstWUE} = P_n/T_i$ (Xu and Zhou, 2008).

**Determination of the plant water status.** Leaves were cut off and their fresh weight was measured. To determine the turgid weight, leaves were kept in distilled water in darkness until they reached a constant weight (full turgor, after 24 h). The relative water content (RWC) was calculated according to the following equation: $\text{RWC} = (\text{fresh weight} - \text{dry weight})/(\text{turgid weight} - \text{dry weight})$.

After leaves were cut off, the leaf water potential ($\Psi_{leaf}$) was measured immediately using a pressure chamber (PMS1000 Instrument; Corvallis, OR). Measurement of the predawn leaf water potential (predawn $\Psi_{leaf}$) began at about 0430 HR and finished before sunrise. In general, midday leaf water potential (midday $\Psi_{leaf}$) was measured between 1230 and 1330 HR. Stem water potential (\(\Psi_{stem}\)) was estimated using the bagged-sealed leaf technique (Zsogon et al., 2015). Briefly, the opposite leaf near a transpiring leaf was covered with aluminum foil and plastic wrap the night before measurement. The pressure potential of the bagged leaf was assumed to provide an estimate of $\Psi_{stem}$.

The evaporative flow method was applied to estimate midday whole-plant hydraulic conductance ($K_{plant}$) using the midday parameters of plant water status (Attia et al., 2015; Simonin et al., 2014; Tsuda and Tyree, 2000), including the whole-plant transpiration rate ($T_{plant-midday}$) and the water potential decrease between soil and leaf ($\Psi_{soil}-\Psi_{leaf-midday}$).

$$K_{plant} = \frac{T_{plant-midday}}{(\Psi_{soil} - \Psi_{leaf-midday})}.$$  

where $\Psi_{soil}$ is the average soil water potential, which was assumed to equal the predawn $\Psi_{leaf}$ because $\Psi_{soil}$ remained relatively constant and came into equilibrium with the canopy water potential (Cole and Pagay, 2015; Jones, 2007). Therefore, midday $\Psi_{soil}$ values were estimated as predawn $\Psi_{soil}$: $T_{plant-midday}$ was determined as the ratio of weight changes to the total leaf area per plant at hourly intervals around midday, which has been described in detail in a previous study (Tsuda and Tyree, 2000).

The crop water stress index (CWSI) was calculated as the difference between the temperatures of leaves and air by (Idso et al., 1981)

$$\text{CWSI} = \frac{dT - dT_{min}}{dT_{max} - dT_{min}},$$

where $dT$ is the difference between canopy temperature ($T_c$) and air temperature ($T_a$), $T_c - T_a$: $dT_{max}$ is the upper limit of the canopy–air temperature difference that can be reached under nonstressed conditions; and $dT_{min}$ is the lower limit of the canopy–air temperature difference under fully stressed conditions. The values for CWSI range from zero to one, where zero indicates no stress and one indicates maximum stress. The upper and

![Fig. 1. Comparison of (A) predawn and (B) midday leaf water potential between low- and high-vapor pressure deficit (VPD) treatments. NS = nonsignificant.](image-url)

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lower baselines of \( \Delta T_{\text{max}} \) and \( \Delta T_{\text{min}} \) were determined according to the relationship of the canopy–air temperature difference \( (T_c - T_a) \) vs. VPD under nonstressed and fully stressed conditions, respectively, as described in detail in a previous study (Zhang et al., 2017). Leaf temperature was determined with a digital infrared thermometer (model GM320; Maizhe Co., Ltd, Shanghai, China) on seven healthy and mature leaves distributed randomly along the different layers of the canopy.

**Determination of plant growth and morphological parameters.** Plant samples were homogeneous for morphological criteria at the beginning of the experiment. The final morphological parameters were determined from the harvest. Plant biomass and leaf area were measured every 20 d after transplanting until fruits were harvested. Leaf area per plant was measured using an Li-3000 leaf area meter (LI-COR Inc.). Samples were dried at 80°C to a constant weight and weighed.

Growth analysis parameters, including the relative growth rate (RGR), net assimilation rate (NAR) and leaf area ratio (LAR), were calculated from the equations (De Groot et al., 2001):

\[
\text{RGR} = \frac{1}{W_1} \frac{\Delta W}{\Delta t} = \frac{\ln(w_2) - \ln(w_1)}{(t_2 - t_1)} \quad [3]
\]

\[
\text{NAR} = \frac{1}{L} \frac{\Delta W}{\Delta t} = \frac{w_2 - w_1}{t_2 - t_1} \times \frac{\ln(L_2) - \ln(L_1)}{L_2 - L_1} \quad [4]
\]

\[
\text{LAR} = \frac{L}{W} = \frac{L_1 + L_2}{2} \quad [5]
\]

where \( W_1 \) and \( W_2 \) are the biomasses of whole plants at times \( t_1 \) and \( t_2 \), and \( L_1 \) and \( L_2 \) are the total leaf areas of whole plants at times \( t_1 \) and \( t_2 \). \( t_1 \) and \( t_2 \) were 20 and 40 d after transplanting in the current study, respectively.

**Transpired water consumption and WUE.** Whole-plant water use efficiency (WUE\(_{\text{plant}}\)) was calculated as the ratio of the shoot biomass to the cumulative amount of water transpired (Kadam et al., 2015). The agronomic WUE (WUE\(_{\text{yield}}\)) was calculated as the ratio of the fruit yield to the cumulative amount of water transpired (Reina-Sánchez et al., 2005).

**Fig. 2.** Effect of vapor pressure deficit (VPD) regulation on the energetic and hydraulic driving forces for water transport along the soil–plant–atmosphere continuum. Solid and dotted lines represent a series of pathways of water flow in liquid and vapor phases. \( \Delta \Psi \) represents water potential drawdown between two compartments of the soil–plant–atmosphere continuum.

**Fig. 3.** Comparison of the typical diurnal variations in (A) plant transpiration and (B) leaf relative water content in the high- and low-vapor pressure deficit (VPD) treatments on a sunny day. Values are the means ± SE (n = 5). Significant differences between high- and low-VPD treatments were compared using Tukey’s test. * Significant at \( P < 0.05 \) or 0.01, respectively.
Results

Energetic and driving forces for water movement. The meteorological data over the whole growing seasons are described in detail in Supplemental Table 1. Regulation of greenhouse VPD had great effects on the energetic and driving forces for water transport. Soil was maintained at a homogeneous water status between the two treatments, as indicated by a nonsignificant difference in predawn $\Psi_{\text{leaf}}$ (Fig. 1A). Midday $\Psi_{\text{leaf}}$ decreased in both the low- and high-VPD treatments, and this depression in midday $\Psi_{\text{leaf}}$ was moderated efficiently by reducing VPD (Fig. 1B). VPD control in relation to both energetic and driving forces for water transport is illustrated in Fig. 2. The driving force at the leaf–air boundary was the greatest along the soil–plant–atmosphere continuum and was reduced substantially in the low-VPD treatment (Fig. 2). Qualitatively, the water driving force at the leaf–atmosphere boundary can be 100-fold larger than that at the soil–leaf boundary.

Plant water status. Regulation of VPD by humidification had great effects on plant–water relations. Diurnal plant transpiration in the two VPD treatments followed similar trends to those of $T_a$ and VPD. The plant transpiration rate decreased substantially in the low-VPD treatment, in which it was reduced by $\approx 35\%$ around midday (Fig. 3A). The diurnal leaf RWC in the two VPD treatments followed similar trends, with dramatic declines around midday (Fig. 3B). Leaf desiccation was moderated effectively by reducing VPD, and the leaf RWC was increased by 5% to 10% around midday (Fig. 3B). Plant hydraulic conductance increased significantly by 28.6% in the low-VPD treatment (Fig. 4A). Plants were less stressed in the low-VPD compartment compared with the high-VPD compartment, according to the value of the midday CWSI (Fig. 4B).

Leaf gas exchange. Reducing VPD increased significantly the photosynthesis rate, $g_s$, and intercellular CO$_2$ concentration (Fig. 5A–C). The leaf transpiration rate and $L_s$ in the low-VPD treatment were reduced substantially by 26.1% and 39.7%, respectively (Fig. 5D and E). Leaf InstWUE was increased substantially by reducing VPD (Fig. 5F).

Plant growth. Dry biomass of leaves was increased by 18.1% by reducing VPD, whereas dry biomass of leaves was unaffected by VPD regulation (Fig. 6A and B). Dry biomass of fruit was increased by 22.5% by reducing VPD.
reducing VPD (Fig. 6C). Overall, reducing VPD increased shoot biomass significantly by 20.5% (Fig. 6D).

Reducing VPD increased substantially RGR and NAR by 23.9% and 79.1%, respectively (Fig. 7A and B). No significant difference in LAR was observed between the two treatments (Fig. 7C).

**Fruit yield, water consumption, and WUE.** The enhancement in plant growth by reducing VPD was accompanied by a substantial improvement in productivity. Reducing VPD increased the fruit yield per plant significantly by 20.1% (Fig. 8A). Cumulative transpired water consumption was decreased significantly by 16.4% (Fig. 8B) in the low-VPD treatments. Along with the improvements in fruit yield and plant biomass, WUE_{yield} and WUE_{biomass} were increased significantly by 43.1% and 40.5%, respectively (Fig. 8C and D).

**Discussion**

Reducing excessive VPD decreased irrigation demand and moderated plant water stress by optimizing driving forces for water transport. Water transport through the soil–plant–atmosphere continuum is a passive process driven by gradients of free energy. The driving force for water transport was determined by water potential gradients between the soil and the atmosphere ($\Delta \Psi = \Psi_{\text{soil}} - \Psi_{\text{air}}$). In natural conditions, soil moisture varies much less than the high-frequency evaporative demand (Caldeira et al., 2014). In addition to its high frequency, $\Psi_{\text{air}}$ reached an excessive negative value at midday in the current study. Consequently, the driving force for water transport was the greatest at the leaf–atmosphere boundary. Qualitatively, the water driving force at the leaf–atmosphere boundary can be 100-fold larger than that at the soil–leaf boundary according to the experimental evidence.

Plant water deficit and physiological disorders occurred under the high VPD condition, despite plants being well irrigated. Experimental evidence demonstrated that evaporative demand generated excessive water driving force and plant water stress. Theoretical analysis in combination with experimental evidence highlighted the implications of VPD control in modulating water transport and improving plant water status. Maintaining optimum ranges of VPD was an efficient solution to reduce excessive atmospheric driving force for water loss, which was confirmed by the experimental evidence. Consequently, reducing the atmospheric driving force led to substantial reductions in the water flow rate and cumulative transpired water consumption in the current study. The transpiration rates of the leaf and the whole plant were reduced by different degrees in the current study, which can be attributed to the great environmental difference between the leaf chamber and greenhouse.

The mass balance between water supply and loss can be expressed as (Martínez-Vilalta et al., 2014; Roddy et al., 2016)

$$\Delta W = J - E = k_{\text{plant}} \times (\Psi_{\text{soil}} - \Psi_{\text{leaf}}) - g_s \times VPD,$$

where $W$ is the water content, $J$ is water uptake and transport, $E$ is the water transpired loss, and $g_s$ is canopy stomatal conductance.

If the mass balance of plant water is to be maintained, $J$ must equal $E$, and then $\Delta W = 0$. Water balance is therefore of fundamental importance to maintain a physiologically
favorable water content and to avoid desiccation. Disruptions in the water balance and plant water stress were the foremost bottlenecks for crop productivity and were moderated efficiently by reducing VPD in the current study.

**Reducing excessive VPD enhanced plant photosynthesis and productivity by improving plant water status.** In addition to reducing water loss, enhancing carbon (biomass) production was also an efficient solution to improving physiologic WUE. Plant productivity processes such as plant growth, biomass, and yield production were improved by VPD regulation. Improvements in biomass and yield production can be attributed to the process of photosynthesis (Zhu et al., 2010). Even a small increase in the photosynthesis rate can translate into large improvements in biomass and yield, because carbon gain is integrated over the growing time and crop canopy (Parry et al., 2011). In the current experiment, the same numbers of nodes and fruit were retained for each plant. Therefore, the enhancement in fruit yield production under low-VPD conditions can be attributed to the increase in individual fruit weight. Growth analysis in combination with gas exchange parameters highlighted the significant role of VPD control in improving plant growth and productivity.

Reducing excessive VPD improved photosynthesis by sustaining stomatal function and improving plant water status. Loss of CO₂ uptake was mitigated efficiently by VPD regulation in the current study. As mentioned, sustained stomatal function for CO₂ uptake in the low-VPD treatment can be attributed to the maintenance of water balance and homeostatic water status. Stomatal closure in response to plant water deficit in vascular plants is believed to be a passive process driven by a reduction in turgor of the guard cells (Buckley, 2005). Because the turgor pressure of stomata is linked directly to plant water status, VPD manipulates the physical and physiological characteristics of guard cells by mediating water transport and affecting leaf hydration. VPD regulation maintains the water balance and relative homeostatic water status by coordinating the vapor loss and liquid supply, as described in Eq. [6]. Combined with a high leaf water potential, the turgor pressure of guard cells was sufficiently high to sustain pore openness for CO₂ use in low-VPD-grown plants.

Comparison of water-productivity improving approaches that regulate atmospheric and soil water status. Water-saving irrigation techniques such as controlled, alternate partial root-zone irrigation and deficit irrigation have been widely recognized as important tools to increase WUE. In our study, WUE improved significantly as a result of a substantial reduction in the loss of transpired water, with no significant reduction in yield production. In the current study, a reduction in transpired water consumption and improvement in plant productivity were achieved simultaneously by regulating atmospheric VPD. WUE based on the criteria of plant biomass and fruit yield was increased by reducing VPD. Considering the trade-off between water loss and carbon gain, the current study suggests that regulating atmospheric VPD had an advantage over water-saving irrigation techniques. This theoretical analysis needs further experimental comparisons between the two techniques.

Although the implication of VPD control in improving WUE was highlighted by theoretical analysis and experimental evidence, VPD regulation faced dilemmas when the theory was applied in agricultural practice. Mechanical application systems such as the pad-fan system or the fogging system were necessary to facilitate VPD control, which required extra investment. In addition to plant-transpired water consumption, the application of fogging for VPD regulation also accounted for agricultural water consumption in greenhouse production. Water savings and WUE improvement over experimental periods will be counterbalanced to some extent by the fogging water. Although low VPD improved cucumber growth and productivity, the negative effect of low VPD was not considered in the current study. Many lines of evidence were provided to indicate that a low VPD reduced the driving force for nutrient uptake and transport, which is especially likely to cause calcium deficiency (Bakker, 1990). In addition, the germination of fungal pathogen spores increases and diseases spread rapidly when VPD is low. Nutritional deficiencies and disease epidemics will affect fruit quality negatively.

The evaluation of environmental control must be based on economic considerations—the best balance between the net profit increase and systems investment. To reduce bottlenecks limiting physiological productivity and economic efficiency, a holistic view of greenhouse environmental regulation is required that incorporates a trade-off between carbon gain and water consumption, yield increase, and fruit quality.

**Conclusions**

Reducing excessive VPD improved the WUE of cucumber by balancing the trade-off between water consumption and plant productivity. Reducing excessive VPD had significant implications for reducing atmospheric water driving forces, which reduced significantly the water flow rate and cumulative
plant transpiration. The reduction in excessive water loss moderated plant water stress efficiently, as leaf hydration, hydraulic limitation, and excessive negative water potential were prevented by maintaining water balance in the low-VPD treatment. The moderation of water stress by reducing VPD sustained stomatal function for photosynthesis and plant growth, which increased fruit yield and shoot biomass substantially. Reduced transpiration in combination with improved plant productivity contributed to a substantial increase in WUE based on the criteria of plant biomass and fruit yield.

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Supplemental Table 1. The average daily meteorological data inside the greenhouse during growing period.

| Environmental parameters | Air temp (°C) | Relative humidity (%) | Vapor pressure deficit (KPa) | Light intensity (Klux) |
|--------------------------|---------------|------------------------|------------------------------|----------------------|
|                          | Control       | Humidification         | Control | Humidification | Control | Humidification | Control | Humidification |
| 16–23 Apr.               | 23.9          | 23.5                   | 66.8    | 78.1           | 1.46    | 0.71           | 13.7    | 13.7           |
| 24 Apr.–1 May            | 25.1          | 24.8                   | 70.1    | 75.6           | 1.37    | 0.87           | 15.6    | 14.9           |
| 2–9 May                  | 26.8          | 25.7                   | 65.5    | 75.1           | 1.56    | 0.89           | 16.9    | 16.4           |
| 10–17 May                | 26.5          | 25.8                   | 63.5    | 72.2           | 1.59    | 0.91           | 17.7    | 17.6           |
| 18–25 May                | 28.7          | 26.5                   | 60.1    | 72.4           | 1.53    | 0.86           | 17.9    | 17.7           |
| 26 May–1 June            | 28.6          | 27.5                   | 60.5    | 75.3           | 1.65    | 0.83           | 19.4    | 19.4           |
| 2–9 June                 | 29.5          | 28.6                   | 56.6    | 70.2           | 1.70    | 0.92           | 19.2    | 19.0           |
| 10–17 June               | 31.8          | 29.9                   | 54.7    | 69.8           | 1.95    | 1.14           | 20.7    | 20.2           |
| 18–25 June               | 33.7          | 31.9                   | 51.9    | 66.4           | 2.14    | 1.37           | 21.5    | 21.5           |
| 26 June–2 July           | 32.6          | 30.2                   | 50.5    | 67.9           | 2.17    | 1.25           | 20.9    | 20.3           |
| 3–10 July                | 33.5          | 31.2                   | 50.3    | 65.4           | 2.26    | 1.18           | 21.2    | 21.0           |
| 11–18 July               | 32.3          | 30.9                   | 52.1    | 66.9           | 2.09    | 1.05           | 20.6    | 20.6           |