Detection of Water Stress in Orchard Trees With a High-Resolution Spectrometer Through Chlorophyll Fluorescence In-Filling of the O_2-A Band

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Abstract—A high spectral resolution spectrometer with 0.065-nm full-width half-maximum was used for collecting spectral measurements in an orchard field under three water stress treatments. The study was part of the FluorMOD project funded by the European Space Agency to develop a leaf-canopy reflectance model to simulate the effects of fluorescence. Water deficit protocols generated a gradient in solar-induced chlorophyll fluorescence emission and tree physiological measures. Diurnal steady-state chlorophyll fluorescence was measured from leaves in the field between June and November 2004 using the PAM-2100 fluorometer to study the effects of water stress on chlorophyll fluorescence. Spectral measurements of downwelling irradiance and upwelling crown radiance were conducted with the narrow-band spectrometer, enabling the canopy reflectance to be obtained at subnanometer spectral resolution and permitting the evaluation of the fluorescence in-filling effects on reflectance in trees under water stress conditions. Diurnal and seasonal measurements showed consistently lower steady-state fluorescence (F_t) and quantum yield ΔF'/Fm' in water-stressed trees, yielding mean values of F_t = 0.38 (well-irrigated) and F_t = 0.21 (water-stressed trees). The agreement between F_t and water potential showed that steady-state fluorescence could be used to detect differences in water stress levels, with determination coefficients ranging between r^2 = 0.48 and r^2 = 0.81 for individual dates. Analysis in the 680–770-nm range showed that the chlorophyll fluorescence in-filling in the O_2-A band at 760 nm is sensitive to diurnal variations of fluorescence and water stress, yielding r^2 = 0.76 (well-watered treatment), r^2 = 0.89 (intermediate stress treatment), and r^2 = 0.7 (extreme stress treatment), demonstrating the close relationships between F_t and in-filling at the crown level.

Index Terms—Chlorophyll fluorescence, fluorescence in-filling, hyperspectral, olive tree, oxygen O_2-A band, remote sensing, water stress.

I. INTRODUCTION

The importance of chlorophyll fluorescence (CF) is being investigated in several studies which focus on the theory, measurement methods, and interpretation, evaluating its relationship with photosynthesis and plant physiological status [1]–[7]. Light-induced oxidative damage is minimized through this protective process, with plant chloroplasts dissipating light energy that exceeds photosynthetic demands [8], [9]. Under increasing stress, plant tissues increase heat production to dissipate excess energy, reducing CF production, at least in the initial and intermediate stages of stress [10]. The relationship between steady-state fluorescence (F_t) and water stress was studied by Flexas and collaborators [11]–[13] showing a strong correlation between F_t and the diurnal variation of stomatal conductance, and to a lesser extent CO_2 assimilation, also affected by irradiance levels and water stress. These results suggested a potential for water stress monitoring in vegetation using solar-induced chlorophyll fluorescence measurements, without requiring active fluorescence sensors.

Several studies attempted to measure the solar-induced chlorophyll fluorescence naturally emitted by plant tissues using passive remote sensing methods, which would enable the global monitoring of plant physiology and photosynthetic functioning at large spatial and temporal scales. As an example, measurements of solar-induced natural fluorescence in vegetation canopies were reported by McFarlane [14] who measured solar-induced fluorescence in a crop canopy using the H-α Fraunhofer line at 656 nm, and Carter [15], [16] using the H-α and O_2-B lines in leaf measurements. Later, Buschmann and Lichtenthaler [17] demonstrated that the solar-induced fluorescence signal is added on top of leaf reflectance signatures, with other qualitative studies suggesting the effect of fluorescence on apparent reflectance could be detected in the red edge spectral region [18]. Later, quantitative assessments in the laboratory of the F_t signal superimposed on the apparent reflectance at both the leaf and canopy levels using time-decay experiments, light blocking methods with long-pass filters in the 400–700-nm region, reflectance difference calculations, and optical indexes demonstrated that F_t contributed up to 2% on the apparent reflectance in the 750-nm spectral region [19]–[21]. Additionally, Campbell [22] using filters to control light induction showed that the contribution of F_t to apparent reflectance in Zea mays (corn) was over 2%.

Recent studies continue to provide additional evidence for fluorescence in-filling of the O_2-A atmospheric oxygen absorb-
tion band at 760 nm, as a detectable feature in the radiance spectra at the near-canopy levels [23]–[26]. At the airborne and far-field scales, Maier and Zarco-Tejada [27], [28] found evidence for the detection of the solar-induced fluorescence signal on apparent reflectance obtained from airborne sensors Reflective Optics System Imaging Spectrometer (ROSIS) and Compact Airborne Spectrographic Imager (CASI) based on the in-filling of fluorescence in the 760-nm atmospheric oxygen absorption band. Despite these initial results, little work has been conducted that quantitatively demonstrates the link between steady-state fluorescence and reflectance under natural light conditions. Although the FRT and SLOPE models were developed to account for the effects of chlorophyll fluorescence on leaf apparent reflectance [19], [29], the need for an integrated leaf-canopy model to account for spectral fluorescence effects on leaf and canopy reflectance initiated the Development of a Vegetation Fluorescence Canopy Model (FluorMOD) funded by the European Space Agency (ESA) (http://www.ias.csic.es/fluormod). Current validation efforts for FluorMOD require the collection of vegetation radiance and reflectance spectra at leaf and canopy levels under different viewing geometries, stress levels, and spectral configurations. As a result of these efforts, the FluorMOD project [30] is making progress on the development of a leaf model that simulates leaf fluorescence, FluorMODleaf [31], and a linked leaf-canopy model that simulated canopy fluorescence, FluorSAIL [32]. The FluorMODgui integrated leaf-canopy model [33] enables the simulation of diurnal effects under different viewing geometries, atmospheric characteristics, and illumination dependency, modeling the effects of natural fluorescence on apparent canopy reflectance.

This manuscript presents progress made as part of FluorMOD validation efforts, assessing whether water stress levels caused by different irrigation treatments would affect the crown-level natural fluorescence emitted, assessing the potential for its detection through the fluorescence in-filling effects in the O$_2$-A band at 760 nm. The study focuses on the detection of steady-state fluorescence superimposed on the canopy reflectance, using water stress as a driver for obtaining different stress and fluorescence levels. Conclusions will facilitate the evaluation of the simulated fluorescence effects on canopy reflectance, assessing the potential for global monitoring of solar-induced chlorophyll fluorescence.

II. MATERIALS AND METHODS

A. Study Site Location and Experimental Design

The field experiment was conducted from June to November 2004 in a 4-ha irrigated olive orchard (Olea europaea L cv. “Arbequino”) located in Córdoba, southern Spain (37.8°N, 4.8°W). The study site is located in an area with Mediterranean climate, with an average annual rainfall of 650 mm concentrated from autumn to spring, and potential evapotranspiration of 1390 mm. The soils in the study area are Typic Xerofluvents, with sandy sliny characteristics, sandy stratum at 1.5-m depth.

The orchard was planted in 1997 in a 3.5 m × 7.0 m grid (408 tree ha$^{-1}$) in a north–south direction. The trees were planted on ridges to avoid flood damage problems because of the drainage and the low percolation rate, using weed killer for soil maintenance. The drip irrigation method used in this study enabled different irrigation treatments over a single experimental area. Fig. 1(a) and (b) shows the study site area, the orchard field and the trees used for this experiment, as imaged by the CASI sensor at 1-m spatial resolution and eight spectral bands in the 400–950-nm spectral range. The CASI sensor flew the study site as part of the AgriSpectra airborne campaign conducted by Spanish and Canadian teams. A chlorophyll fluorescence gradient was sought through three irrigation treatments randomly applied in six rows of 18 olive trees covering an area of 2646 m$^2$, generating three water stress levels: 1) irrigation at 2.8 mm/day (well watered treatment, used as control, R); 2) 0.7 mm/day (deficit treatment, stress treatment S1); and 3) an intermittent treatment, with 1.2 mm/day from June 14, 2004 to July 5, 2004 and from September 6, 2004 to October 19, 2004, with no irrigation from July 5 to September 6 (deficit treatment, stress treatment S2) [Fig. 1(c)].

Trees located in the center of each treatment block were selected for weekly monitoring of tree physiological status, with measurements made of leaf water potential, stomatal conductance, photosynthesis and leaf chlorophyll fluorescence. In addition, crown measurements of reflectance in the 680–770-nm spectral range were conducted on trees under different stress levels and for different viewing geometries.

B. Field Data Collection

1) Leaf Measurements: Weekly measurements were conducted on 11 trees located in the center of the irrigation blocks to avoid treatment edge effects, to monitor water potential ($\Psi_w$) on four trees per treatment for S1 and S2 treatments, and three trees for the control treatment R. Among them, two trees per treatment were selected for weekly measurements of chlorophyll fluorescence, stomatal conductance, and photosynthesis. A Scholander pressure bomb (PWSC Model 3000, Soilmoisture Equipment Corp., California) was used to measure leaf water potential from 11 trees covering the three irrigation treatments, with measurements taken weekly at 10:00 solar time from two shaded leaves per tree located close to the trunk. Stomatal conductance was measured weekly with a leaf steady-state porometer (model PMR-4, PP Systems, Hitchin Herts, Great Britain), every hour from 7:30 to 11:30 solar time from three trees, on one tree per irrigation treatment, on five labeled sun-exposed leaves per tree. Leaf photosynthesis was measured weekly with a CIRAS-1 instrument (PP Systems, Hitchin Herts, Great Britain) at 7:00 and at 10:00 solar time from six trees on two labeled sun-exposed leaves per tree.

Leaf chlorophyll fluorescence measurements were conducted under field conditions using the Pulse-Amplitude-Modulated Fluorometer PAM-2100 (Heinz Walz GMBH, Effeltrich, Germany), an instrument that has been used widely in basic and applied fluorescence research [34]. The instrument enables the monitoring of steady-state and dark-adapted fluorescence features, providing flexibility for scientific research on chlorophyll fluorescence [11]. Therefore, the focus of this validation experiment was on measuring steady-state chlorophyll fluorescence as a function of stress condition, enabling the assessment of natural fluorescence effects on canopy spectral reflectance for remote sensing detection under natural sun light conditions. Procedures
used for measuring steady-state (Ft) and fluorescence quantum yield (ΔF/Fm') were based on standard methodologies as documented in the PAM-2100 manual (Heinz-Walz-GmbH, 1993).

The leaf was positioned in the PAM-2100 leaf clip holder, which exposes a sample area approximately 1-cm in diameter to the fiberoptic light emitter and detector array. Steady-state fluorescence Ft feature was measured along with the effective quantum yield ΔF/Fm', which denotes the actual efficiency of PSII photon capture in the light by closed Photosystem II (PSII) reaction centers, determined as $\Delta F/Fm' = (Fm' - Ft)/Fm'$, where Fm' is the maximal fluorescence of a preilluminated sample with PSII centers closed, and Ft is the fluorescence at
steady-state [35], [36]. An average of 50 exposed leaves per tree were used for $F_t$ and $\Delta E/Fm'$ measurements five times per day between 6:30 h and 11:30 h solar time during June and November 2004.

2) Canopy Spectral Measurements: Apparent reflectance data were obtained for the top of tree crowns using a 0.065-nm full-width half-maximum (FWHM) Ocean Optics HR2000 fiber-optics spectrometer (Ocean Optics, Dunedin, FL) installed on a 7.0-m height pole. The sensor head included a cosine collector for downwelling irradiance and a bare fiber for upwelling radiance (Fig. 2). The 0.065-nm FWHM HR2000 spectrometer provided spectral measurements in the 680–770-nm range with 2048 channels from top of the crowns in a diurnal data collection series coincident with leaf fluorescence measurements. The spectrometer was specifically designed for the experiment, and built with grating H11 of 1800 lines/mm, a 5-µm slit, an L2 detector collection lens, an OF1-OG590 longpass filter, and a set of high-reflectivity AgPlus mirrors model SAG-UPGD-HR to enhance sensitivity of the instrument for high-resolution measurements in low-light level experiments (all these model parts belong to the instrument manufacturer, Ocean Optics, Dunedin, FL). The instrument was connected to a 10-m long, 600-µm optical fiber using a CC-3 VIS-NIR cosine corrector-diffuser (Ocean Optics, Dunedin, FL). Due to dark current sensitivity of the instrument as function of ambient temperature, a Peltier thermally insulated box Model PT-100 (Magapor, Zaragoza, Spain) was used to house the spectrometer, keeping the internal temperature fixed at 24 °C ± 1 °C during field measurements.

Irradiance ($E_0'$) calibration of the spectrometer attached to the fiber with the cosine corrector-diffuser was conducted in the laboratory using the LS-1-CAL calibrated tungsten halogen NIST-traceable light source (Ocean Optics, Dunedin, FL). Radiance (L) calibration was conducted with a Spectralon panel (Labsphere, North Sutton), calculating the apparent reflectance (R) of the tree crown with the equation $R = \pi \cdot L/E_0'$, which includes the reflected radiance and the fluorescence emission effects. Wavelength calibration of the instrument was conducted with the Hg-Ar HG-1 light source (Ocean Optics, Dunedin, FL). Aerosol optical measurements were acquired with a Microtops II sunphotometer (Solar Light, Philadelphia) measuring at 440-, 500-, 675-, 870-, and 936-nm spectral bands. The sunphotometer was connected to a GPS model GPS-12 (Garmin, Olathe, KS) for simultaneous readings of geographic location, altitude and solar geometry at the time of spectral acquisitions.
The spectral measurements were made over selected tree crowns under the three different water stress treatments, measuring irradiance and crown radiance at high spectral detail inside the O₂-A (760.5 nm) oxygen absorption band to study the detection of natural fluorescence in-filling [23]–[26] on spectral reflectance as function of stress status and diurnal variation of chlorophyll fluorescence. The prominent irradiance absorption lines due to atmospheric oxygen can be seen in Fig. 3(a) (O₂-A and O₂-B bands) measured with the HR2000 spectrometer at 0.065-nm FWHM in the 680–780-nm spectral range, observing the O₂-A absorption band in detail [Fig. 3(b)]. Internal structure and frequency features due to molecular oxygen can be observed in the very high spectral resolution of less than 1 Å FWHM used in this experiment. Field-measured vegetation radiance collected from pure tree crowns under sun conditions shows the oxygen absorption bands [Fig. 3(c)] and the spectral detail inside the O₂-A band in the radiance signal [Fig. 3(d)]. Calculation of the apparent reflectance spectra using the measured downwelling spectral irradiance [Fig. 3(a)] and vegetation radiance spectra [Fig. 3(c)] generates a peak or in-filling at the oxygen absorption maxima shown on the red edge vegetation spectra [Figs. 3(e) and (f)] which has been shown with potential to quantify natural chlorophyll fluorescence emission from vegetation. The high spectral resolution used in this experiment facilitates the observation of the structure of the in-filling of the O₂-A band, assessing the potential effects of the FWHM of the instrument used on the detection of the fluorescence signal superimposed on the canopy reflectance. The absolute in-filling amplitude was calculated using a band affected by fluorescence emission in O₂-A (R₇₅₀₋₅₉) and another band outside the fluorescence in-filling (R₇₇₅₋₅₉), obtaining a measure of the in-filling absolute amount by the subtraction R₇₅₀₋₅₉ − R₇₇₅₋₅₉. The shape of the red edge reflectance spectrum collected with the HR2000 spectrometer, showing the in-filling at the O₂-A band [Fig. 3(e)] indicates the characteristic vegetation spectral shape in the red edge region normally acquired with other spectrometers of lower spectral resolution, suggesting that proper methods for spectral data collection were used. Crown spectral measurements and in-filling calculations were conducted in diurnal trials from trees under each water stress level, acquiring spectra and fluorescence measurements every 30 min between 8:30 and 11:30 solar time. Sample reflectance spectra obtained with this high-resolution spectrometer from the crown of two trees using the large pole are shown in Fig. 4 noting the red edge features between 700 and 750 nm, and at higher spectral detail between 755 and 765 nm with the figure insert showing the in-filling at 760 nm due to the vegetation natural fluorescence emission.

III. RESULTS

Leaf water potential measurements conducted at 10:00 solar time from the 11 trees monitored over the entire summer showed large variations in water stress as function of the irrigation treatment [Fig. 5(a)]. As expected, trees subjected to stress treatments S1 and S2 with lower irrigation doses were more stressed than control R trees with irrigation for full evapotranspiration (ET) [Fig. 5(b)]. Trees with maximum water stress (S1) reached −3.3 MPa on September 30, before the first rainfalls in early October. Control irrigation (R) trees showed stable water potential over the course of the experiment, reaching average values of −1 MPa with a minimum of −0.6 MPa, whereas the intermittent deficit irrigation treatment S2, yielded water potential values in between the R and S1 irrigation treatments [Fig. 5(b)]. A full recovery of the water potential at the end of the experiment was achieved as a consequence of the rainfalls after the summer. Consistently, both stomatal conductance [Fig. 5(c) and
Large differences were found between the extreme irrigation treatments R and S1 for both Ft and ΔF/Φm, demonstrating that effects of deficit irrigation and water stress on steady-state fluorescence measures over the season were detected, tracking the water potential recovery at the end of the experiment. Results of the diurnal measurements for Ft [Fig. 8(a) and (b)] and ΔF/Φm' [Fig. 8(c) and (d)] on two different dates with increasing water stress condition [Fig. 8(a) and (c) for August 5th, \( \Psi_w = -1 \) MPa (R), \(-1.7\) MPa (S2), \(-1.8\) MPa (S1)] and [Fig. 8(b) and (d) for August 26th, \( \Psi_w = -1.3\) MPa (R), \(-2\) MPa (S2), \(-2.4\) MPa (S1)] indicated that a maximum value for Ft was found between 7:30 and 8:30 solar time, decreasing afterwards. Moreover, the diurnal studies showed that the largest differences between treatments were found at 10:00 solar time for both Ft and ΔF/Φm', with small differences in the emitted fluorescence signal as function of water stress at earlier times of the day. The diurnal measurements showed consistent lower Ft and ΔF/Φm' values in water stressed trees, yielding mean values of Ft = 0.28 in well-irrigated trees with \( \Psi_w = -1.3\) MPa (R) as compared to Ft = 0.21 in stressed trees with \( \Psi_w = -2.4\) MPa (S1) on August 26th. Diurnal variations of steady-state fluorescence suggested that midday values acquired at 10:00 solar time for Ft and ΔF/Φm' were therefore better for separating different irrigation treatments as function of water stress status.

(d)] and photosynthesis rates [Fig. 5(e) and (f)] were greater for the low-stressed well-watered treatment (R) trees at the two times of measurements in the morning (8.00 solar time) and midday (11.00 solar time). Stomatal conductance measured at 11:00 solar time was affected over the course of the experiment as function of stress status, increasing from 329 mmol/m² at the time of maximum stress to 680 mmol/m² at recovery for treatment R, from 203 to 573 mmol/m² for treatment S2, and from 80 to 507 mmol/m² for treatment S1. The photosynthesis rate measured at 10:00 solar time throughout the experiment consisted of a reduction for treatment S1, as illustrated by 10.35 \( \mu \)mol/m² for well-irrigated trees and 6.5 \( \mu \)mol/m² for S1 stressed trees at the time of maximum stress (September 27). These measurements of water potential, conductance, and photosynthesis suggest that the experimental design to create a gradient in stress condition as function of water deficiency was successful, therefore enabling the study of the imprint of chlorophyll fluorescence on apparent crown spectral reflectance.

Diurnal variations of stomatal conductance acquired throughout the season between 7:30 and 11:30 solar time also showed a consistent behavior (Fig. 6) yielding maximum conductance rates for the control trees at 8:30 solar time, with a general decrease in the trend afterwards. Diurnal rates showed consistent trends for the control R [Fig. 6(a)] as compared with the deficit treatment S2 [Fig. 6(b)] and the highest stress S1 [Fig. 6(c)], yielding lower conductance rates over the course of the day. Both the seasonal and diurnal variation of conductance and photosynthesis measures, which showed a recovery at the end of the season, enabled the study of fluorescence effects as function of stress status. The leaf chlorophyll fluorescence measurements collected from selected tree crowns diurnally and throughout the season showed that steady-state fluorescence Ft [Fig. 7(a) and (b)] and fluorescence quantum yield ΔF/Φm' [Fig. 7(c) and (d)] for the low-stressed trees (R) were higher than for the high water-stressed treatments (S2, S1) at 8:00 [Fig. 7(a) and (c)] and 10:30 solar time [Fig. 7(b) and (d)] with measurements in three trees per irrigation treatment. Large differences were found between the extreme irrigation treatments R and S1 for both Ft and ΔF/Φm', demonstrating that effects of deficit irrigation and water stress on steady-state fluorescence measures over the season were detected, tracking the water potential recovery at the end of the experiment. Results of the diurnal measurements for Ft [Fig. 8(a) and (b)] and ΔF/Φm' [Fig. 8(c) and (d)] on two different dates with increasing water stress condition [Fig. 8(a) and (c) for August 5th, \( \Psi_w = -1 \) MPa (R), \(-1.7\) MPa (S2), \(-1.8\) MPa (S1); Fig. 8(b) and (d) for August 26th, \( \Psi_w = -1.3\) MPa (R), \(-2\) MPa (S2), \(-2.4\) MPa (S1)] indicated that a maximum value for Ft was found between 7:30 and 8:30 solar time, decreasing afterwards. Moreover, the diurnal studies showed that the largest differences between treatments were found at 10:00 solar time for both Ft and ΔF/Φm', with small differences in the emitted fluorescence signal as function of water stress at earlier times of the day. The diurnal measurements showed consistent lower Ft and ΔF/Φm' values in water stressed trees, yielding mean values of Ft = 0.28 in well-irrigated trees with \( \Psi_w = -1.3\) MPa (R) as compared to Ft = 0.21 in stressed trees with \( \Psi_w = -2.4\) MPa (S1) on August 26th. Diurnal variations of steady-state fluorescence suggested that midday values acquired at 10:00 solar time for Ft and ΔF/Φm' were therefore better for separating different irrigation treatments as function of water stress status.
The relationships obtained between leaf water potential $\Psi_x$ and fluorescence measures $F_t$ and $\Delta F/Fm'$ for single days and for the entire experiment throughout the season indicated that steady-state fluorescence measures are good indicators of water stress status at the tree level (Fig. 9). The agreement between $F_t$ and $\Psi_x$ on different dates throughout the season for which coincident water potential and fluorescence measurements were made, showed that steady-state fluorescence $F_t$ could detect differences in water potential, with $r^2 = 0.48$ [July 21; Fig. 9(a)], $r^2 = 0.8$ [August 4, Fig. 9(b)], $r^2 = 0.7$ [August 25, Fig. 9(c)], and $r^2 = 0.81$ [September 1, Fig. 9(d)]. For the entire experiment, the agreement between leaf water potential and $F_t$ [$r^2 = 0.54$, Fig. 9(e)] and $\Delta F/Fm'$ [$r^2 = 0.37$, Fig. 9(f)] suggested that steady-state fluorescence $F_t$ can potentially be used as a consistent indicator for water stress in olive tree crops.

The diurnal measurements conducted with the Ocean Optics HR2000 spectrometer at 0.065-nm FWHM in the 680–770-nm range demonstrated that the fluorescence in-filling at the O$_2$-A band was detected and contributed to the tree crown reflectance signal, manifested in a sudden increase of the apparent reflectance at 760 nm (Figs. 3, 4, and 10). The amplitude of the 760-nm peak, calculated as $R_{700.59} - R_{759.5}$, potentially associated with the emission of natural fluorescence, was compared diurnally with steady-state fluorescence $F_t$ measurements collected at the same time from the trees under the different stress levels (Fig. 10). The results obtained in diurnal trials between $F_t$ and the O$_2$-A peak amplitude (Fig. 11) yielded correlations of $r^2 = 0.76$ (for a tree with $\Psi_x = -0.825$ MPa, treatment R well irrigated), $r^2 = 0.80$ ($\Psi_x = -1.05$ MPa, stress treatment S2) and $r^2 = 0.7$ ($\Psi_x = -3.35$ MPa, stress treatment S1), indicating the close relationships between $F_t$ and the fluorescence in-filling signal at the crown level on individual trees. These results suggest the potential monitoring of diurnal changes in fluorescence through apparent reflectance spectra at the canopy level. Moreover, results for the diurnal measurements on all trees under the three different stress conditions yielded $r^2 = 0.61$ (Fig. 12), showing that well-irrigated trees (R) exhibited higher $F_t$ and in-filling values than stressed S1 and S2 trees. These results suggest that variations in natural fluorescence emission were successfully tracked through inferred reflectance spectra on all trees under different water stress conditions, enabling the identification of stress levels. Nevertheless, the dependency of the emission peak on reflectance bidirectional reflectance distribution function (Fig. 10) still requires critical attention due to the known changes as a function of the viewing geometry and solar angle that accompany diurnal changes [32].

**IV. Conclusion**

Apparent reflectance measurements obtained from tree crowns with a high-resolution Ocean Optics HR2000 spectrometer at 0.065-nm FWHM in the 680–770-nm range demonstrated that the observed fluorescence in-filling in the O$_2$-A band at 760.5 nm varied with water stress status in orchard trees.
Diurnal and seasonal measurements acquired over the course of the summer experiment showed consistently lower Ft and $\Delta F/Fm'$ values in water-stressed trees, yielding mean values of $Ft = 0.38$ in well-irrigated trees with $\Psi_x = -1.3$ MPa (R) and $Ft = 0.21$ in stressed trees with $\Psi_x = -2.4$ MPa (S1). Diurnal measurements of steady-state fluorescence demonstrated that midday values acquired at 10:00 solar time for Ft and $\Delta F/Fm'$ were better for separating different irrigation treatments as function of water stress status.

The relationships obtained for leaf water potential with Ft and $\Delta F/Fm'$ for single days and for the full dataset comprising the entire experiment showed that steady-state fluorescence Ft is a good indicator of water stress status at the tree level in olive canopies. The agreement between Ft and $\Psi_x$ throughout the season showed that steady-state fluorescence could detect differences in water potential levels, with determination coefficients ranging between $r^2 = 0.48$ and $r^2 = 0.81$ for individual dates. For the entire experiment, the agreement between leaf water potential and Ft ($r^2 = 0.54$) and $\Delta F/Fm'$ ($r^2 = 0.37$) demonstrated that steady-state fluorescence Ft shows potential as a consistent indicator of water stress in olive tree crops.

The amplitude of the 760-nm peak, calculated as $R_{700,530} = R_{750,530}$, associated with the emission of natural fluorescence, was compared diurnally with steady-state fluorescence Ft measurements collected at the same time from the trees under different stress levels. The results obtained in diurnal trials between Ft and the O2-A peak amplitude yielded $r^2 = 0.76$ (for a tree with $\Psi_x = -0.825$ MPa, well-irrigated treatment R), $r^2 = 0.89$ (R2 = -1.05 MPa, stress treatment S2) and $r^2 = 0.7$ ($\Psi_x = -3.35$ MPa, stress treatment S1), demonstrating the link between Ft and in-filling of the 760-nm apparent reflectance at the crown level on individual trees. Results for the diurnal measurements considering all trees under study yielded $r^2 = 0.61$, suggesting that natural fluorescence was successfully monitored through reflectance spectra on all trees under different water stress conditions.

These results obtained at canopy level for water stress monitoring through fluorescence detection are in agreement with studies conducted by Mora [37] at the leaf level, and with canopy results by Moya [23], showing that fluorescence emission can be detected at the leaf level and on a corn canopy under diuron herbicide penetration using the O2-A band. In this case the study was conducted on orchard trees, showing that water stress variations were responsible for corresponding fluorescence effects on canopy reflectance, enabling its detection with a narrow-band spectrometer under natural field conditions. Conclusions may lead toward important implications for water stress monitoring and water scheduling applications in agriculture, suggesting the potential application of high spectral resolution reflectance data for monitoring natural chlorophyll fluorescence emission at the canopy level as an indicator of water stress.

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