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UV-LED irradiation for biofouling reduction in drip irrigation emitters fed with wastewater effluent†

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Crop irrigation with treated wastewater effluent using drip irrigation has become common as the demand for water supply has increased. Because of the quality characteristics of treated wastewater and the narrow and winding geometry of the drip emitter's structure, it is susceptible to clogging. Emitter clogging reduces flow and increases flow variability between emitters that can lead to water stress on crops, thereby reducing crop yield. Several methods to minimize emitter clogging have been suggested and applied; however many drawbacks are associated with them. The use of UV-LEDs (UV light-emitting diodes) is a non-chemical disinfection method that holds great promise for disinfection and biofouling prevention in irrigation systems. In this research, biofouling formation potential was investigated for 12 weeks, in a large pilot-scale irrigation rig consisting of three parallel pipelines, comparing three disinfection treatments: UV-LED, chlorine, and no treatment. The results indicate that the discharges of UV-LED and chlorine-treated lines were similar. However, analyzing the internal fouling material of the opened drippers revealed the significant advantage of the UV-LED treatment, when both OCT (optical coherence tomography) image processing and EPS (extracellular polymeric substance) secretion within the clogging substances indicated significant biofilm inhibition by UV-LED irradiation as compared to the other alternatives. The present study is a proof-of-concept of a new approach of using UV-LED irradiation for minimizing biofouling formation in emitters fed with treated wastewater. UV-LED technology has great potential to become an attractive and feasible alternative for replacing chlorine as a water disinfection technology, specifically for agriculture use.

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Water impact

The use of UV-LEDs, a non-chemical disinfection method, has great potential for disinfection and for limiting biofouling formation in irrigation systems, and may replace chlorine for agriculture use. This research provides evidence on the efficiency of biofilm inhibition by UV-LED irradiation compared to chlorine or no treatment disinfection, at a pilot-scale drip irrigation rig fed with treated wastewater.

1. Introduction

Agriculture irrigation using treated wastewater effluent (TWW) has become common in the last decades as the demand for water supply has increased, especially in arid and semi-arid areas.^{1,2} Drip irrigation combined with using TWW has many positive aspects, from advancing water sustainability to energy saving and improving agricultural yields.³ However, due to the emitters' narrow and winding

internal structure they may clog, particularly when TWW is used, due to increased chemical and biological fouling. Emitter fouling may reduce flow and the uniformity of water application, reducing the overall irrigation water volume and increasing the variation between the drippers, respectively.^{4–9}

Several methods to minimize and/or eliminate emitter clogging have been suggested and used, including pre-filtration, which reduces the amount of suspended organic and inorganic matter entering the drip irrigation system. However, filtration cannot remove all the suspended matter and eventually sediments and microbiological growth accumulate in the emitters.^{10,11} Another possible method is pressure flushing, in which the pipelines are rinsed by increasing the hydraulic shear force within the pipe system. A flushing velocity of 0.6 m s^{−1} was reported to be adequate; however it must be performed often to dislodge and transport accumulated sediment.^{6,12} Acidification was

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‡ In memory of Barak White who sadly passed away in May 2022, before completing his research.



reported as a method to prevent chemical precipitation, yet it may adversely affect soil pH.^{13,14} The most common method for reducing emitter bio-clogging by bacteria and algae is chlorination.^{4,6,13,15} That said, there are no unequivocal and unified guidelines for the required chlorine concentrations, exposure duration and timing, for controlling drip-irrigation bio-clogging.^{4,16} In addition, irrigation with chlorine-containing effluent has its drawbacks: chlorine may damage the dripper structure, leading to a significant decrease in dripper performance;⁵ the effluent may damage the plants' roots;¹⁵ when the effluent contains high ammonia concentrations, chlorination will result in the formation of chloramines which last longer in the distribution system but have lower-efficiency as biocides; organic matter in the effluent increases chlorine demand necessitating the addition of a higher chlorine dose to achieve the same efficiency, and may yield trihalomethanes (THMs) and other disinfection-by-products (DBPs), exhibiting negative health effects.¹⁷ Finally, chlorination entails relatively high capital and operational costs due to the need to transport, store and dose chlorine in the field. Therefore, a cheaper, simpler, and more environmentally-, health- and crop-friendly method for emitter biofouling reduction is warranted.

Ultraviolet (UV) irradiation has been used as a common and effective technology for water disinfection and reduction of biofouling in water systems and membranes.^{18,19} UV disinfection has several advantages over conventional chemical disinfection, such as no chemical addition, storage, and dosing. It does not remain in the water and no harmful DBPs are formed and it has little effect on the environment. However, conventional UV disinfection utilizes mercury lamps, which have disadvantages such as large size and high energy demand. In addition, the lamps are fragile and contain toxic mercury, which is hazardous to the environment and requires proper disposal.²⁰ Recently, UV light-emitting diodes (UV-LEDs) have emerged as a new source for UV irradiation to replace mercury vapor lamps, and have been investigated and used as a tertiary wastewater treatment.^{20–24} UV-LED technology has several advantages over conventional mercury UV lamps, such as diversity of radiating wavelengths (from deep UV to near UV regions; 210–360 nm), small point light source, mercury-free, durability, compactness, extremely short warm-up time (intermittent-flow friendly), allowing versatile integration, lower energy consumption, and longer lifetime.^{20,25,26} Since disinfection depends on the spectral sensitivity of the target microorganism rather than the UV source, the efficacy of LEDs for disinfection of protozoa, viruses, and bacteria was proven to be at least as effective as low-pressure UV lamps for a given wavelength and dose combination.^{27–34} Bacteria (including biofilm-formers) and protozoa such as *Cryptosporidium* and *Giardia* that pose a health risk are easily inactivated at low UV doses, while virus inactivation requires higher doses.²⁷ The UV-LED wavelength output is tunable based on material composition,²⁵ making it an ideal technology for optimization of treatment objectives, as the response of each microorganism to different wavelengths is unique.^{32,33} Few studies have

already evaluated the efficacy of UV-LEDs for treated wastewater; however, these are limited;^{22,23,32,35} in particular, most of them have focused on small-scale systems and there is a lack of case studies involving flow-through UV-LED reactors in large pilot-scale systems of treated wastewater for agricultural use as drip irrigation emitters. In addition, to the best of our knowledge, no attempt has been made to evaluate biofouling clogging inside drippers fed with UV-LED-disinfected effluent at a full-scale pilot drip irrigation rig in long-term studies (few months).

Several methods for analyzing biofouling clogging in drip irrigation have been reported, some of them require extraction and destruction of the clogging sample. Others are non-destructive, proposing to study the samples without extracting them.⁸ Monitoring emitter discharge is a direct method, which is non-destructive and is suitable for both laboratory-scale and field studies. Dripper flow rate and flow uniformity among drippers are affected by biofilm formation and therefore allow the evaluation of clogging rate and severity.⁸ Extraction methods, indicative of biofilm formation and quantification, include analysis of biofilms and bacterial activity using extracellular polymeric substances (EPSs). EPS quantification has been widely and efficiently used for studying clogging in drip irrigation.^{8,36} Quantifying biofilm formation can also be studied by optical coherence tomography (OCT), which provides a three-dimensional view of the sample, and allows the identification of regions most susceptible to biofilm clogging.^{7,9,37}

The main objective of the work was to study the effect of UV-LED irradiation on biofouling formation and to quantify fouling formation mechanisms in emitters fed with TWW. In this study we analyzed the fundamental principles of UV-LED disinfection and evaluated its inactivation efficacy for biofilm-forming bacteria using a collimated-beam apparatus and a flow-through reactor at the laboratory, and at a full-scale pilot drip irrigation rig fed with TWW. To understand biofouling formation in drippers at a full-scale pilot drip irrigation rig we used non-destructive direct methods based on accurate-automatic discharge monitoring and extraction and destruction methods based on OCT and EPS characterization.

2. Materials and methods

2.1 Target organisms and microbiological procedures

Pure culture of green fluorescent protein (GFP) vegetative *Bacillus subtilis* (NCIB 3610) was used as the representative microorganism and a biosimulator for the disinfection studies, and as the target microorganism for inactivation tests at the laboratory. *B. subtilis* is a non-pathogenic Gram-positive bacterium that is highly tractable. Consequently, *B. subtilis* frequently serves as a model organism for the study of Gram-positive bacteria in several applications, amongst them studying biofilm formation.³⁸

The spiking solution was prepared by placing a single colony of roughly 1–2 mm in diameter in a sterile Erlenmeyer



flask containing 20 mL nutrient broth. The Erlenmeyer flask was shaken for 18 h at 100 rpm and 30 °C. The final spiking stock solution contained $\sim 10^{10}$ CFU per 100 mL. *B. subtilis* concentration was determined using the filter membrane method; samples were seeded on nutrient agar (HiMedia, India) Petri dishes and incubated for 24 h at 30 °C and colonies were counted.

2.2 Experimental setups

2.2.1 UV-LED collimated beam apparatus. Bench-scale collimated beam experiments were conducted following Sholtes and Linden (2019) using a UV-LED device (PearlLab Beam, AquiSense Technologies, USA; Fig. 1a and b),³⁹ consisting of three different nominal wavelengths: 255, 265, and 285 nm. In this study, the 285 nm LED lamp was used, as this was the wavelength of the flow-through UV-LED reactor. UV irradiation at 285 nm is disinfection-wise less efficient than 255 nm as the 285 nm wavelength is less absorbed by nucleic acids compared to 255 nm. However, the fluence of the 285 nm UV-LED is about 10 times larger (data not shown) and its serviceable life is much longer (AquiSense Technologies, personal communication), making it more suitable for the studied application.

UV irradiance was measured using a spectrometer (USB2000 + UV-VIS-ES, Ocean Optics, USA) at different distances. The spectrometer probe was fastened below the lamp to ensure precise measurement at the center of the UV-LED lamp, while the collimated beam was adjusted to measure at a 6 cm diameter radial template, at 10° and 5 mm intervals Fig. 1c. Absolute irradiance was measured and used for calculating the Petri factor.⁴⁰

Average UV fluence obtained on the surface of the irradiated solution sample was measured with iodide-iodate chemical actinometry, based on Rahn *et al.* (2003).⁴¹ The actinometry solution was photolyzed into triiodide, the concentration of which was directly related to the radiation dose the solution had experienced. Aliquots of 10 mL of

iodide solution were placed in a 6 cm Petri dish, 9 cm from the UV-LED source and then irradiated between 10 s and 3 min with continuous stirring. The UV-LED irradiation fluence was calculated using 0.37, iodide-iodate quantum yield at 285 nm,⁴² and the intensity was calculated according to Rahn *et al.* (2003).⁴¹

2.2.2 Flow-through UV-LED reactor. A flow-through UV-LED reactor (47 mL total internal volume) emitting at 285 nm was used (PearlAqua Micro, AquiSense Technologies, USA; Fig. 1d). The fluid enters the reactor and flows through two chambers: a cooling chamber (21 mL) followed by an irradiation chamber (26 mL). According to the manufacturer, the LEDs' operational lifetime is 8000 h. The reactor unit has an input voltage of 12 V DC and requires a minimum flow rate of 0.1 L min⁻¹.

The examined flow-through UV-LED reactor is patent protected, and its internal geometry and structure were unknown to us. Therefore, indirect approaches for characterizing the UV-LED flow-through reactor were required, using three methods: tracer experiments, chemical actinometry and biodosimetry (bacteria inactivation experiments).

Tracer experiments were conducted using the flow-through UV-LED reactor to understand and quantify flow regime elements based on Friedler and Gilboa (2010).⁴³ Briefly, the NaCl solution served as a tracer, and its flow rate, varying between 0.1 and 0.6 L min⁻¹, was controlled by a step feed peristaltic pump. The solution electrical conductivity was recorded at 1 s intervals at the outlet of the reactor (Metrohm 912 Conductometer, Switzerland). The flow regime was analyzed using time distribution curves, and the reactor hydraulic efficiency was estimated by flow characterization parameters (obtained from the distribution curves): average residence time (t_{ave}), the time of peak tracer concentration appearance (modal; t_p), and the time when 10% of the final concentration reached the outlet (t_{10}). The ratios between these parameters and the theoretical residence time (T – reactor volume divided by the flow rate) were calculated (for more details, see Friedler and Gilboa, 2010 (ref. 43)). The flow regime

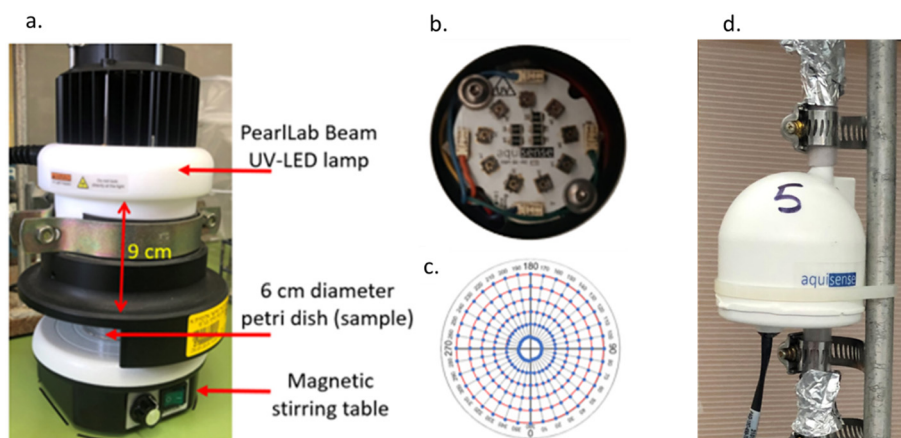


Fig. 1 UV-LEDs: a) collimated beam setup; b) top view of the lamps; c) radial template for measuring irradiation intensity distribution; d) flow-through reactor.



in the UV-LED reactor was further assessed using the Rebhun-Argaman model (1965) which calculates the proportions of the plug flow, dead spaces, and degree of mixing.⁴⁴

In the actinometry experiments, prior to feeding the actinometric solution, the UV-LED reactor was rinsed with distilled water and the examined flow rates were adjusted. The feed was then instantly switched to the iodide-iodate solution. For calculating the irradiance intensity of the UV-LED reactor, the iodide-iodate solution was fed at flow rates identical to the ones used in the tracer experiments (0.1–0.6 L min⁻¹). As aforementioned, the inner structure of the PearlAqua reactor is patent protected and unknown to us, and specifically the area exposed to the radiation (cm²) is not explicitly defined. Therefore, to quantify UV irradiation intensity, an assumption had to be made, and the area exposed to the radiation was calculated by multiplying the irradiation chamber volume by the absorption ratio of the iodide-iodate solution at 285 nm (1 cm⁻¹).

Biodosimetry was used for evaluating the UV-LED efficacy applied to the water using the flow-through UV-LED reactor.⁴⁵ In biodosimetry, the unknown UV dose of the flow-through reactor is compared to a well-known UV dose of the collimated beam (as fundamentally characterized in section 2.2.1). The dose-response curve (*i.e.*, log inactivation of the examined bacterium *vs.* UV dose), obtained using the collimated beam, serves as a standard curve for back-calculation of the UV dose of the UV-LED at the flow-through reactor at each flow rate. The latter is defined as the RED (reduction equivalent dose) of the flow-through reactor. The RED was later adjusted for uncertainties and biases to produce the validated dose of the reactor for the operating conditions examined.

Two types of solutions were examined by biodosimetry: phosphate buffered saline (PBS) and synthetic treated wastewater (STWW; for the procedure, see ESI† data, S1) with a transmittance of 95 and 80%, respectively. 20 L of each water matrix was prepared daily and spiked with the target bacteria (*B. subtilis*) to achieve an initial concentration of ~10⁶ CFU per 100 mL (measured at the beginning of the experiment). The water matrices were then exposed to UV irradiation in the UV-LED flow-through reactor or the collimated beam, and the RED of the flow-through reactor was calculated from the dose-response of the collimated beam. At the collimated beam, a water matrix (PBS or STWW) of 10 mL was placed in a 6 cm Petri dish, 9 cm from the UV-LED source and then irradiated between 10 s and 30 min with continuous stirring. For the UV-LED flow-through reactor experiments, 20 L of the examined water matrices were pumped through the reactor at flow rates of 0.1–2.0 L min⁻¹. The *B. subtilis* concentration (inflow and outflow) was measured on nutrient agar using the filter membrane method (method 9215 (ref. 46)).

2.2.3 Full-scale pilot drip irrigation rig. A pilot-scale drip irrigation rig was constructed in the laboratory to comparatively study the effectiveness of UV-LEDs and chlorine in reducing biofilm formation, clogging and the

performance of the emitters. The experiments used STWW as the irrigation water that was fed into 3 drip-irrigation lines, each receiving a specified treatment: chlorination, UV-LED, and control (no treatment; Fig. 2a and b).

STWW was prepared daily in a 500 L tank, stirred, and spiked with *B. subtilis* at a concentration of ~10⁶ CFU per 100 mL. Mixed STWW was then pumped (PV55 50 Hz, Pedrollo, Italy) through a 130 µm disk filter (Arkal 1", Amiad, Israel), and the pressure was reduced to 1.4 atm (¾" PRV 1.4 atm, Bermad, Israel) and diverted to the three treatment lines. Each line consisted of a 25 m long 16 mm (external) diameter tube (to ensure the 30 min residence time needed for the chlorination treatment, see below) followed by a drip line with 10 integral pressure compensating 1 L h⁻¹ drippers (UniRam™ AS, Netafim, Israel), spaced 0.2 m apart. The UniRam drippers contain an internal diaphragm that constantly vibrates according to the pressure changes to prevent sedimentation of organic matter and other pollutants (self-cleaning). Irrigation was performed 5 d per week for 6 h d⁻¹ and was operated for 12 weeks, representing the irrigation cycle of a typical field crop.

At the head of each line, a disinfection treatment was executed, either chlorination or UV-LED irradiation. The third line was not disinfected (control). Chlorination was achieved by pumping hypochlorite (as sodium hypochlorite) with an injection pump (gamma per L, ProMinent, USA) at the treatment line head ensuring 30 min contact time (see above). To ensure residual chlorine of 0.5–1.0 mg L⁻¹ as Cl₂, chlorine demand was measured using the DPD colorimetric method, according to APHA (2017).⁴⁶ At the head of the UV-LED line, 2 flow-through UV-LED reactors were placed in series and were wired to a computer for internal heat measurements to ensure that the reactors do not overheat.

Chlorination was performed as a proactive measure, which was conducted when the relative emitter discharge along any of the two treatment lines dropped below 80% (residual chlorine at the dripper outlet of 0.5–1 mg L⁻¹ as Cl₂ for 3 and 2 h for chlorine and UV-LED lines, respectively). Further, once a week the flow-through UV-LED reactors were detached from the treatment line and washed with freshwater to reduce the potential of biofouling inside the reactors.

The *B. subtilis* (the target bacteria) concentration was measured twice a week at the three drippers' outlet of each line and was compared to its initial concentration measured at the STWW tank. Finally, log inactivation was calculated (log (*B. subtilis* concentration at the tank/*B. subtilis* at the outlet of the dripper)).

2.3 Biofouling clogging analysis

For biofouling clogging analysis at the full-scale pilot drip irrigation rig, we used non-destructive methods, which were continuously employed for 12 weeks, in parallel to extraction and destruction methods, performed at the end of the experiment. To assess which treatment provides better dripper functionality, statistical analyses were performed for



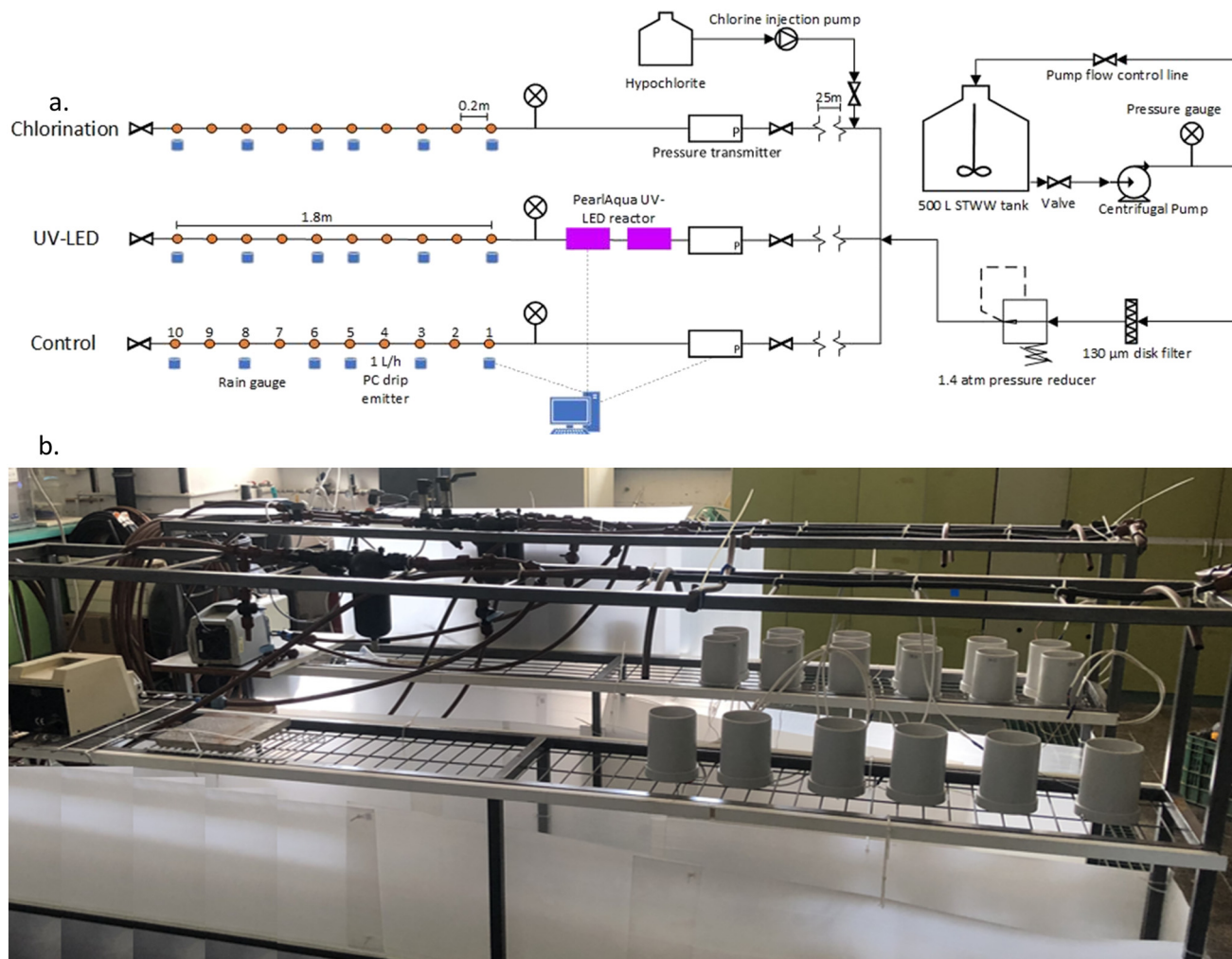


Fig. 2 a. Schematic of the laboratory pilot-scale drip irrigation rig; b. the laboratory pilot-scale drip irrigation rig.

all non-destructive (flow rate analysis; section 2.3.1) and destructive methods (chemical and microbial analysis and extracellular polymeric substances; section 2.3.2), using Student's *t*-test and ANOVA (analysis of variance), calculated using the statistical software JMP Pro.

2.3.1 Flow rate measurement and analysis (non-destructive method). Flow rate monitoring is a direct-non-destructive method of clogging assessment within drippers. Flow rates of selected drippers (1st, 3rd, 5th, 6th, 8th, and 10th dripper) from each line were continuously measured by placing tipping-bucket rain gauges (Monsoon, Agrolan, Israel) underneath, collecting the drippers' emission. Measurements were recorded using a computer collecting the tipping count and timestamp of the rain gauge inner swing (5 ml). The tipping-bucket rain gauges were first calibrated by measuring the exact volume passing through one movement of the inner swing to convert swing tipping to volume of water. The flow rate data, measured from each treatment line using the rain gauges, were recorded using a computer and then normalized to the initial flow rate values measured at the beginning of the experiment using distilled water:

$$NQ_i^t = \frac{q_i^t}{q_i^0} \quad (1)$$

NQ_i^t Normalized flow rate of dripper *i* at time *t* [%]

q_i^t Flow rate of dripper *i* at time *t* using STWW [L h⁻¹]

q_i^0 Initial flow rate of dripper *i* using distilled water (*t* = 0) [L h⁻¹].

The relative dripper discharge along a treatment drip line was calculated as follows:

$$\text{Dra} [\%] = \frac{100}{n} \cdot \sum_{i=1}^n NQ_i^t \quad (2)$$

Dra Relative emitter discharge along the drip line [%]

n Number of drippers measuring the flow rate.

The uniformity of drip emitter flow rate presents a second key indicator to the drip line flow efficiency:

$$\text{UC} [\%] = 100 \cdot \left(1 - \frac{1}{nNQ} \sum_{i=1}^n |NQ_i^t - \overline{NQ}^t| \right) \quad (3)$$



$$\overline{NQ}^t = \frac{1}{n} \sum_{i=1}^n NQ_i^t \quad (4)$$

UC Uniformity coefficient [%]

\overline{NQ}^t Average normalized flow rate at time t [$L\ h^{-1}$].

2.3.2 Dropper analysis by extraction and destruction methods. Extraction and destruction methods, indicative of biofilm formation and quantification were performed at the end of the pilot-scale experiment (after 12 weeks). The drippers were cut open and the internal fouling material was analyzed using the following methods: traditional chemical and microbial analysis, EPS and OCT (5 drippers for chemical analysis, 2 drippers for microbial analysis and 3 drippers for EPS and OCT).

Chemical analysis. The chemical analysis procedure included cutting open the drippers, placing the drippers, including the internal diaphragm, in 20 mL glass vials with distilled water, ultrasonication for 15 min at 37 kHz and 100% strength (DC-250H digital ultrasonic cleaner, MRC, Israel) pouring solution through 1.2 μm glass-fiber filters (GFA, Whatman, UK) and washing the drippers by airbrush spraying distilled water. Drippers were filtered in pairs to ensure enough fouling material for measurement. Filtered rinse water was used for measuring total suspended solids (TSS) and volatile suspended solids (VSS),⁴⁶ the filtrate was analyzed for TOC and TN using a TOC-TN analyzer (TOC-V CPH, equipped with a total nitrogen measuring unit, TNM-1; Shimadzu, Japan), positive ions (Mg^{2+} , Na^+ , Fe^{3+} , K^+ , Ca^{2+}) using a high-resolution array ICP emission spectrometer (PLASMAQUANT® PQ 9000 ELITE; Analytik Jena AG, Germany), and negative ions (F^- , Cl^- , NO_2^- , Br^- , NO_3^- , PO_4^{3-} , SO_4^{2-}) using an 881 Compact IC pro-Anion-MCS with chemical suppression by the Metrohm Suppressor Module, equipped with an 836 Compact Autosampler (Metrohm AG, Switzerland). Anion separation was performed on a Shodex column IC-SI-52 4E (4.0 mmID \times 250 mmL). A 3.6 mM sodium carbonate (Na_2CO_3) solution was applied as an eluent.

Microbial analysis. Microbial analysis was achieved by rinsing the cut open drippers (including the internal diaphragm) with 20 mL PBS into sterile 50 mL Falcon tubes. The tubes were then vortexed and shaken (TS-400 orbital shaker, MRC, Israel) for 0.5 h to break/release the fouling structure. Using the filter membrane method, samples were seeded as mentioned in section 2.1.

Extracellular polymeric substances (EPS). The dropper inner side (including the internal diaphragm) was rinsed with 20 mL of PBS and the solute was collected in a tube. Tubes were vortexed and shaken. The EPS protocol was based on the Liu and Fang (2002) method which included biofilm extraction using the addition of sodium chloride (0.9%) and sodium carbonate (0.5%) for breaking the EPS bonds, followed by heating (0.5 h, 80 $^{\circ}C$) and centrifugation (20 min at 6000 rpm; Biofuge Primo Heraeus Centrifuge; Kendro Laboratory Products, Germany).⁴⁷ The supernatant extracted substance was filtered through a 0.45 μm filter (Merck Millipore,

Germany) and dialyzed through a membrane with a molecular weight cutoff of 3.5 kDa (SnakeSkin®; Thermo Fisher Scientific, Inc., USA) to remove undesired ions and retain the EPS. The EPS protein and carbohydrate were separately assessed by following the procedure of two assay kits (Pierce™ BCA Protein Assay Kit, Thermo Fisher Scientific, Inc., USA and total carbohydrate assay kit, Abcam, Zotal, Israel, respectively) and measured using a plate reader (infinite 200Pro, Tecan Trading, Switzerland) at 562 nm and 490 nm, respectively.

Optical coherence tomography (OCT). Optical coherence tomography (OCT) is a noninvasive imaging technique enabling high-resolution, cross-sectional imaging of a wide range of highly scattering media, such as biological tissues. OCT was used to study the biofilm structure inside the drippers of the three treatment lines (UV-LED, chlorine, and control), at the end of the experimental period (12 weeks). Dropper system valves were closed at both ends to keep the drippers wet and then the dropper system was disconnected from the irrigation line. The inside regions of interest of the cut-open drippers were scanned and snapshotted in 2D and 3D images using OCT apparatus (Thorlabs GmbH, Dachau, Germany) combined with high-performance data acquisition software (Software ThorImage OCT 4.4). The winding internal channel was divided into six sections (Fig. 3) to study biofouling in different parts of the channels. Three drippers were cut-open for each treatment line, each area was snapshotted three times and the obtained images were analyzed by image processing analysis. Based on Lequette *et al.* (2021), the OCT images were converted into the 8-bit gray-scale⁶ and an image processing code was developed to detect pixels associated with the plastic tube of the emitter and pixels associated with the biofilm clogging using MATLAB R2021b (MathWorks®, version 2021b). The total intensity of pixel matrix was calculated and signified the biofilm thickness. The data of each position (x , y) were binarized and the developed script was used for estimating the ratio of pixels associated with biofilm clogging to the total image pixels, meaning the biofilm percentage obtained for each treatment.

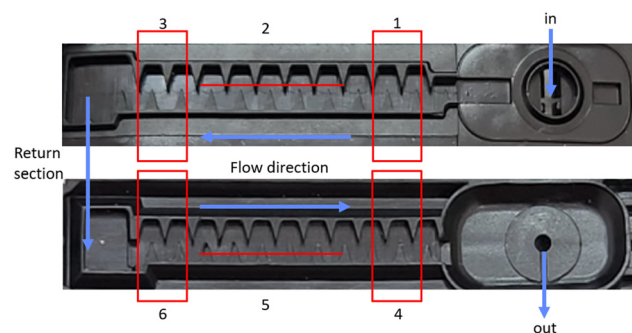


Fig. 3 Internal image of the geometric channel of the drippers and the six sections scanned by OCT. Blue arrows indicate the flow direction. Areas 1,3,4 and 6 were 3D scanned and areas 2 and 5 were 2D scanned.



3. Results and discussion

3.1 UV-LED collimated beam characteristics

The irradiance of the UV-LED collimated beam was quantified by spectrophotometry and actinometry. The spectral emission measured by spectrophotometry revealed that the peak wavelength was slightly different from the nominal one given by the manufacturer (283.1 nm *vs.* 285 nm), and the full width at half-maximum was about 15 nm. The maximum irradiance flux, 9 cm from the UV-LED source, measured at the center of the water sample, was $50.2 \mu\text{W cm}^{-2}$ and the total irradiance flux was 1.9 mW cm^{-2} . The total irradiance flux observed by actinometry was 0.55 mW cm^{-2} , 3.45 times lower than the value obtained by the spectrophotometry method (the same phenomenon was observed for 255 and 265 nm irradiance, data not shown). The irradiance uniformity of the collimated beam (Fig. 4) yielded a Petri factor of 0.835 (6 cm dish at 9 cm distance from the UV-LED source), which is slightly lower than reported ones (~ 0.9).^{24,40} Bolton and Linden (2003) stated that the Petri factor is affected by the design of the collimated beam apparatus, where high petri factors are obtained when the collimated beam apparatus is well designed.⁴⁰ Moreover, if the light source was further from the reactor, the petri factor would likely improve.

3.2 Characterization of the flow-through UV-LED reactor

As stated above, to determine the residence time distribution and the flow regime in the UV-LED reactor a step-feed injection was applied at various theoretical residence times. Flow characterization parameters were determined by using the cumulative residence time distribution function and the residence time distribution function obtained from the tracer experiment, for each examined flow rate. t_{ave}/T and t_p/T ratios, for all flow rates, were higher than 1.0, indicating that the flow distribution in the reactor was not uniform and that there were dead spaces. In addition, the t_{10}/T ratio increased with the flow

rate from 0.4 (flow rate: 0.1 L min^{-1}) to 0.7 (0.6 L min^{-1}), meaning that the first 10% of the tracer reached the outlet after 40–70% of the theoretical time. The results suggest that at the examined flow rates, some mixing occurred in the reactor. The use of Rebhun and Argaman's model confirmed this assumption since the model generated 60–70% degree of perfect mixing of the effective reactor volume.

The UV-LED flow-through reactor radiation intensity was calculated experimentally by actinometry for all examined flow rates and the lamp intensity was calculated to be $5.1 \pm 1.2 \text{ mW cm}^{-2}$. Thus, the UV-LED flow-through reactor radiation intensity was one order of magnitude higher than the irradiance flux of the collimated beam, due to the internal structure of the flow-through reactor and number of LEDs within it. Each LED is an individual source of radiation; hence the number of LEDs and their position inside the reactor affect the total intensity.

The RED of the flow-through reactor was calculated by biosimetry using the dose-response curve obtained from the thoroughly characterized collimated beam (section 3.1; Fig. 5a). The RED curve of the UV-LED reactor matches expectations, with higher inactivation for greater doses and lower flow rates (longer exposure time Fig. 5b and c). The UV transmittance (UVT) of the examined solutions affected the results of dose-response curves and the RED per flow rate. STWW exhibited a lower dose-response curve and RED per flow rate due to absorption and shading effects (UVT $\sim 80\%$) resulting from the presence of dissolved and particulate matter, as compared with the PBS solution (UVT $\sim 95\%$), which was clear from the dissolved and particulate matter.

3.3 Full-scale pilot drip irrigation rig

As mentioned, the pilot-scale drip irrigation rig was constructed to comparatively study the effectiveness of three treatments (UV-LEDs, chlorine, and control; no treatment) in the reduction of biofilm formation using direct non-destructive methods (flow rate measurement and analysis) and extraction and destruction of the clogging samples.

Characterizing the flow-through UV-LED reactor provided data for determining the required UV-LED dose in the full-scale pilot drip irrigation rig. The “UV-LED-treated” line contained 10 integral pressure compensating drippers of 1.0 L h^{-1} flow rate each, meaning that the UV-LED reactor disinfected a total flow of 10 L h^{-1} (0.167 L min^{-1}). According to Fig. 5, the UV-LED reduction equivalent dose (RED) delivered to STWW (UVT $\sim 80\%$) by adjusting the flow-through UV-LED reactor to 0.167 L min^{-1} was 17 mJ cm^{-2} , based on a 2.5 log inactivation. Connecting two UV-LED reactors in series would result in an additive dose of about 34 mJ cm^{-2} .

3.3.1 Flow rate measurement and analysis (a non-destructive method). As aforementioned, the flow rate dripping through the emitters was continuously measured by 6 tipping-bucket rain gauges, for each line, and recorded using a computer (Fig. S1†). The relative drip emitter

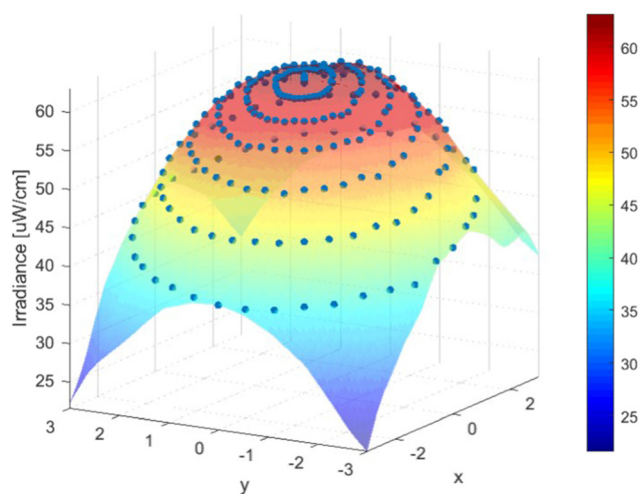


Fig. 4 Irradiance profile of UV-LED radiation of the batch collimated beam (285 nm, measured 9 cm from the UV-LED source).



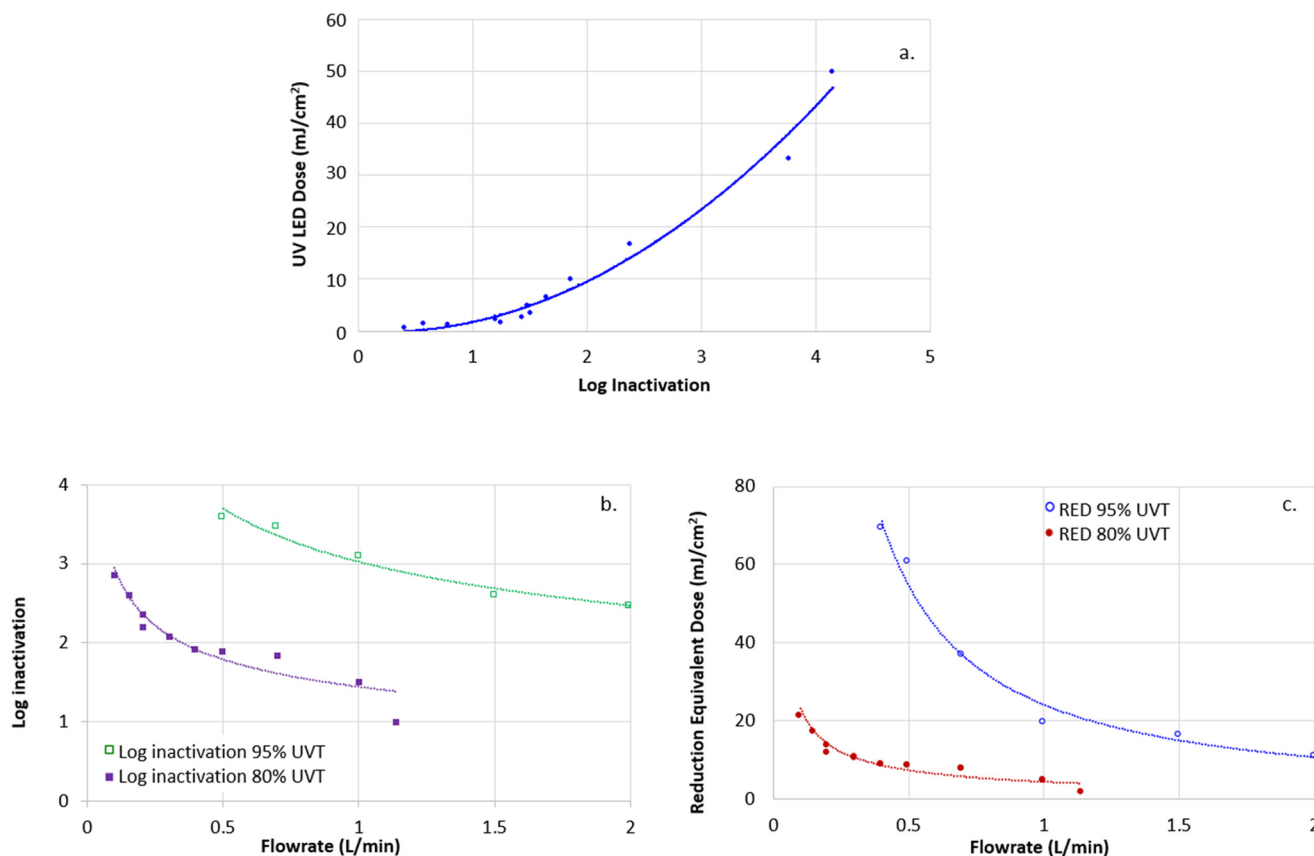


Fig. 5 RED calculated by biosimetry. a. UV-LED dose response curve for the STWW solution (UVT 80%): log inactivation vs. UV-LED dose at the collimated beam apparatus; b. log inactivation of the UV-LED flow-through reactor in two different solutions (UV transmittance); c. RED curves of the UV-LED flow-through reactor in two different solutions (UV transmittance).

discharge along the control line (no treatment) remarkably decreased after 12 weeks of operation, decreasing to 60–70% of the initial discharge, while the discharge of the chlorine and UV-LED lines remained quite high (90–100% and 80–90% of the original discharge, respectively; Fig. 6a). Thus, the

drip line flow efficiency remained high for the UV-LED and chlorine lines and decreased for the control line, implying clogging development in the control line. In total, proactive chlorination was performed 9 and 3 times in the chlorine and UV-LED lines, respectively. It is worth noting that not

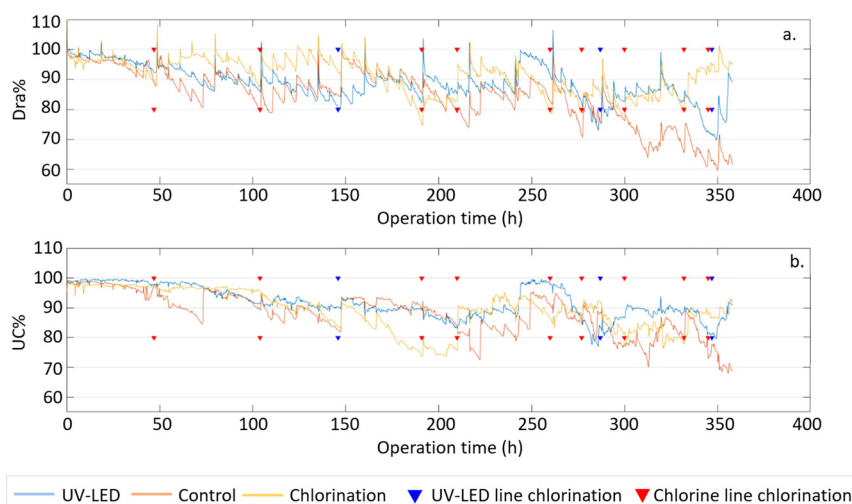


Fig. 6 Analysis of flow through the dripper lines: a) relative overall drip emitter discharge along the line (Dra) vs. operation time. b) Uniformity coefficient (UC) vs. operation time. Triangles – maintenance of the pipelines by proactive chlorination.



only the proactive chlorination in the chlorine line was more frequent and aggressive (3 vs. 2 h) but also the frequency of chlorination increased with the progress of the experiment. This hints that biofilm formation was higher in this line than in the UV-LED line (see also Fig. 8).

Differences between the three treatments were proven to be statistically significant (ANOVA, $p < 0.05$, measured throughout the study), meaning that although a small difference between chlorine and UV-LED lines was observed, it was statistically significant. Similar results (also statistically significant) were obtained for the uniformity coefficient (UC; Fig. 6b) with the UC of the control line being lower (70%) than the UC of chlorine and UV-LED lines (>90%).

Flow rate measurement and temporal recording help to understand the clogging development along the dripper lines. This estimates the presence of clogging and does not require extraction of samples and thus the destruction of drippers, and therefore can be used continuously in the field. However, flow measurement provides an indirect estimation and does not necessarily reflect the early stages of clogging as extraction methods may be able to do.

3.3.2 Biofilm analysis using extraction and destruction methods. After 12 weeks (360 irrigation hours), drippers (10 drippers per line) were cut open and the internal fouling material was analyzed. TSS, VSS, TOC and TN exhibited the

same trend, being lower in the biofilm of the chlorine and UV-LED lines than in the biofilm of the control line (Fig. 7, top). However, a statistically significant difference was found only for VSS, when the VSS concentration at the chlorine and UV-LED lines reduced by 40% compared to the control line. Reduction in the TOC concentration between the control and the UV-LED lines was also statistically significant. It was not surprising that these two parameters were significantly affected as they are indicators of organic matter, such as biofilm-forming bacteria, while TSS and TN are not specific indicators for organic matter and are less affected by the disinfection treatments. Comparing chlorine and UV-LED lines indicated a reduction of about 30% in the TOC concentration at the UV-LED line, however, this difference was not found to be statistically significant.

No clear trend was observed in the concentrations of the examined positive and negative ions at the tree lines (data not shown).

The *B. subtilis* concentration at the biofilm of the UV-LED line was 3 orders of magnitude lower than in the biofilm of the chlorine and control lines (Fig. 7, bottom). Interestingly, the chemical chlorination effect on the *B. subtilis* concentration was not significant. Following chlorination, the *B. subtilis* concentration in the clogging

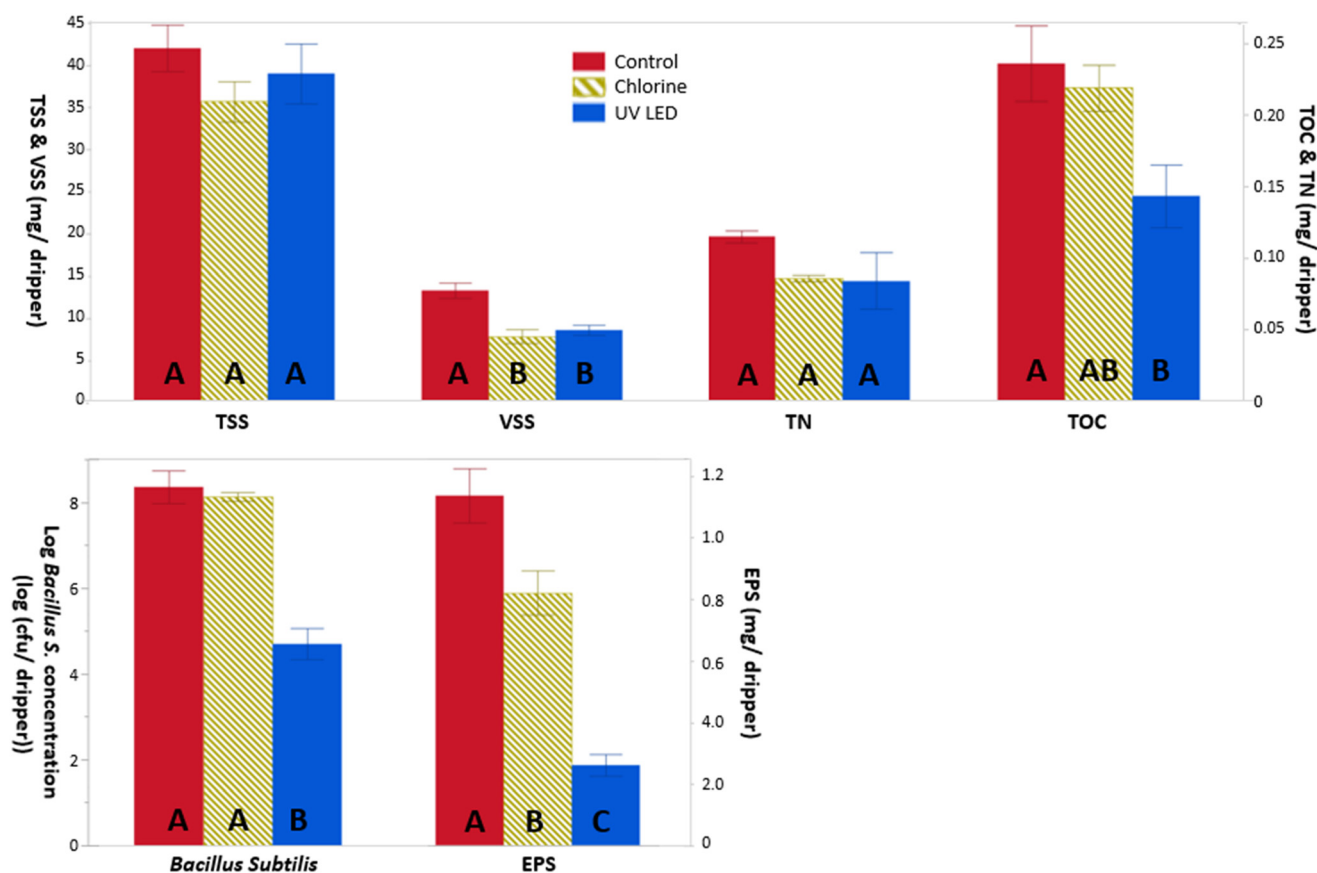


Fig. 7 Chemical (top) and biological (bottom) biofilm analysis extracted from the three dripper lines. Different letters signify significantly different averages ($p < 0.05$).



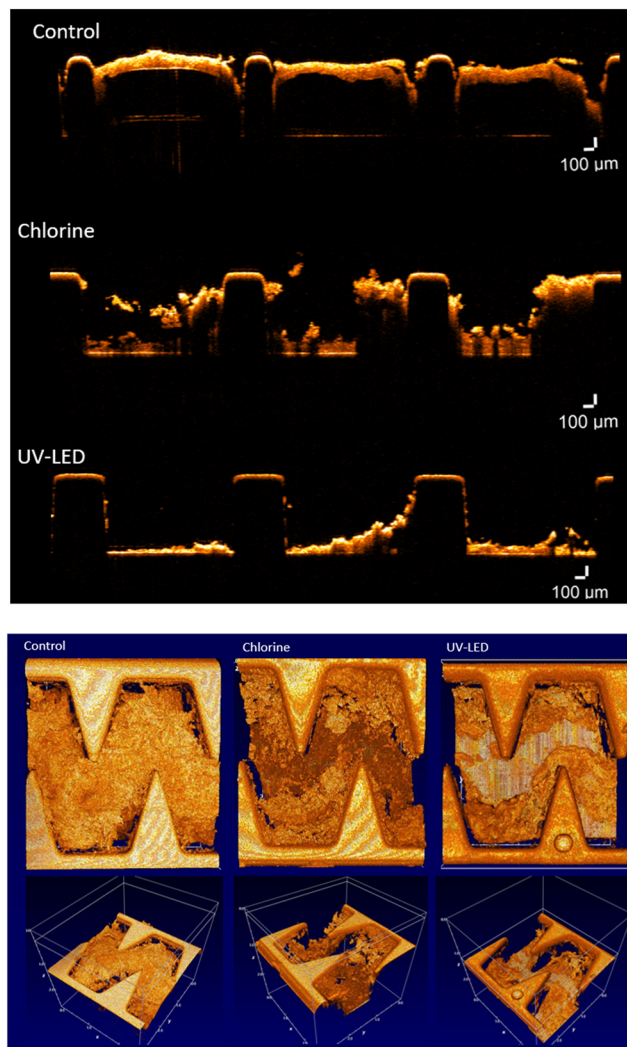


Fig. 8 2D (top) and 3D (bottom) OCT images for all three examined treatments.

substances was similar to the concentration measured within the emitters of the control line. Hence, the *B. subtilis* concentration was much more affected by UV-LED disinfection than by chlorination.

EPSs, which are essential to ensure the maintenance and stability of the biofilm on any surface, such as emitters, were affected by the three different treatments (Fig. 7, bottom). The disinfection treatments (chlorination and UV-LED) could inhibit the secretion of EPSs within the clogging substances. The content of EPSs in the clogging substances was slightly lower for the chlorine emitter line compared to the control line, with a reduction of 30% (t -test, $p < 0.01$). These results are in line with Chen *et al.* (2022) who studied the effects of disinfection methods, including chlorine, on microbial EPS production and on membrane fouling.⁴⁸ The authors reported that EPSs might benefit disinfection residual bacteria which are typically biofilm-forming bacteria, such as *B. subtilis*, by adhering the bacteria to the solid surface, leading to bacteria regrowth and EPS accumulation on the

surface of the system. Increasing chlorine dose might have improved these results, as stated by Song *et al.* (2017), who indicated that the EPS content gradually decreased as chlorination and chlorine duration increased.⁴ However, high chlorine doses might damage the dripper internal structure.⁵

The impact of the UV-LED line on EPSs within the clogging substances was significantly higher, generating 80% and 68% reduction (t -test, $p < 0.01$) in the EPS clogging substances compared to the control and chlorine lines, respectively, thereby substantially minimizing the risk of emitter clogging. It should be noted that in this study we used synthetic treated wastewater to reduce unknown parameters. Real treated wastewater effluent would contain significant amounts of dissolved organic matter, including EPSs that were not simulated with the examined synthetic effluent.

3.3.3 Biofilm analysis using OCT. Fig. 8 reveals the biofilm development in the winding internal channel of the drippers after 360 operation hours, and describes the biofilms' morphology and their dimensional architecture, as obtained in the three examined treatment lines. The intensity of the bright color indicates biofilm accumulation on the emitter structure (black color). The brighter the color, the thicker the biofilm, and thus the higher the ability of biofilms to induce clogging. It is evident from the figure that both treatments (UV-LED and chlorine disinfection) reduced the amount of biofilm accumulated within the dripper.

An in-house image processing code was developed to quantitatively estimate the biofilm thickness and the proportion of the emitter cross sectional area covered by biofilm. The proportion of biofilm at the chlorine and control line were found to be similar, 43%, while at the UV-LED line the percentage of biofilm cover was lower, 32%. The biofilm thickness (estimated by the intensity function of the gray-scale pixels) was 7% and 37% lower at the chlorine and UV-LED lines compared to the control line, respectively, meaning that the chlorine dose used in this study was insufficient to reduce the level of clogging. Higher doses might improve the chlorination performance. On the other hand, the percentage of biofilm cover and biofilm thickness was significantly influenced by the UV-LED disinfection, where OCT images show that both the biofouling thickness and percentage of biofilm cover decreased, as compared to the control line.

4. Conclusions

To improve the efficiency of drip irrigation emitters fed with treated wastewater, it is desired to minimize biofouling clogging. In the present study, the potential of using UV-LED irradiation as a treatment to decrease biofouling formation was investigated as a measure to inactivate biofilm-forming bacteria present in treated wastewater effluent. To demonstrate the fundamental principles of UV-LED disinfection and to evaluate the UV-LED inactivation efficacy, bench-scale experiments were conducted, and UV-LED irradiation was characterized using a collimated-beam



apparatus and a flow-through UV-LED reactor. Using the characterization data obtained by the bench-scale experiments, a full-scale pilot drip irrigation rig imitating a drip irrigation emitter line was designed and constructed, where biofilm formation was investigated.

The pilot-scale irrigation rig was fed with STWW and consisted of three parallel drip-irrigation lines, each for specific treatment: UV-LEDs, chlorine, and control (no treatment), where the effectiveness to reduce biofilm formation due to the three treatments was compared. The direct non-destructive method of accurate-automatic discharge monitoring at a single dripper scale indicated that the relative drip emitter discharge and the uniformity coefficient along the control line significantly decreased after 12 weeks of operation, while the discharge and uniformity coefficient of the chlorine and UV-LED lines remained quite high, meaning that the drip line flow efficiency remained high for the UV-LED and chlorine lines and decreased for the control line.

After 12 weeks, the drippers were cut open and the biofilm developed within the examined pipeline drippers was analyzed. Concentrations of the chemical parameters indicate that the biofilms were lower in the chlorine and the UV-LED line drippers than in the control emitters. Microbial analysis revealed however that only the UV-LED treatment affected the biofilm development, with *B. subtilis* concentrations at the chlorine and control lines being similar and much higher than the *B. subtilis* concentration at the UV-LED line. EPS secretion within the clogging substances was inhibited by both disinfection treatments (chlorination and UV-LED irradiation) compared to the control line, with the UV-LED disinfection having a significant advantage, which had a meaningful impact on the EPS within the clogging substances. Image processing of the OCT analysis supports these results; a lower percentage of biofilm cover and lower biofilm thickness was obtained at the UV-LED line compared to the chlorine and control, meaning that high clogging potential was observed for the control and chlorine lines and limited clogging at the UV-LED line, though a higher chlorine dose might improve its performance.

The overall results indicate that UV-LEDs can play a critical role in reducing biofouling of drip irrigation fed with treated effluent compared to no treatment and chlorination. UV-LEDs may be advantageous further in that they can be located along the irrigation line as an integral part of the system and do not require any chemical addition, they are more environmentally friendly and are likely to reduce operation complexity (making a disinfectant storage tank and dosing equipment redundant), and therefore may serve as a suitable replacement to chlorine. However, there is a lack of information regarding the economic feasibility of using UV-LED irradiation as a treatment to decrease biofouling in emitters, such as initial capital investment, operation cost, energy demand, etc. Thus, future studies should focus on economic aspects of UV-LED irradiation for agriculture use.

The present study is a proof-of-concept of a new approach of using UV-LED irradiation for limiting biofouling formation in emitters fed with treated wastewater. The emerging UV-LED technology has great potential to become an attractive and feasible alternative to chlorine disinfection, and specifically, for minimizing biofouling in emitters fed with treated wastewater.

Data availability

Data for this article, including all experimental results, are available on OneDrive-Technion.

Conflicts of interest

There are no conflicts to declare.

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