



Supplement of

Impact of meteorological conditions on the biogenic volatile organic compound (BVOC) emission rate from eastern Mediterranean vegetation under drought

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S1. Preparation of calibration curve for GC–MS

The calibration curve evaluation was conducted with standards of selected volatile organic compounds (VOCs), including monoterpenes (MTs) and sesquiterpenes (SQTs). Two calibration series were conducted. For each series, different solutions were injected into the same measurement sorbent tubes. Then the linear regression fitting function was evaluated based on calculation of the peak area counts vs. VOC mass (μ g). The first-series calibration was applied for six VOCs, where the sampled solution base was mixed with 5 μ L of each compound. Among them, *cis*- β -ocimene and β -caryophyllene were detected in our research. In the second series of calibrations, we calibrated α -humulene, germacrene D, and α -farnesene. Some details of the standards for the VOCs are shown in Table S1.

Table S1. Summary of calibration compounds' input.

Compound name	Density (g mL ⁻¹)	Standard volume added (µL)
<i>cis</i> -β-Ocimene	0.8	5
β-Caryophyllene	0.905	5
α-Humulene	0.88	5
Germacrene D	0.85	5
α-Farnesene	0.81	5

From the initial solution, we prepared 12 and 7 solutions with different concentrations as summarized in Tables S2 and S3 for the first and second calibration series, respectively.

Solution	Volume of solutes	Methanol solvent (µL)	Concentration (ng µL ⁻¹)	Mass of standard in 4 µL injected in the tube (ng)
GIL-080221	5 μL	500	~10,000	
Sol 1	100 µL of GIL-080221	900	1000	
Sol 2	250 μL of Sol 1	750	250	1000
Sol 3	500 µL of Sol 2	500	125	500
Sol 4	500 µL of Sol 3	500	62.5	250
Sol 5	500 µL of Sol 4	500	31.25	125
Sol 6	500 µL of Sol 5	500	15.625	62.5
Sol 7	500 µL of Sol 6	500	7.8125	31.25
Sol 8	480 µL of Sol 7	520	3.75	15
Sol 9	500 µL of Sol 8	500	1.875	7.5
Sol 10	500 µL of Sol 9	500	0.9375	3.75
Sol 11	533 µL of Sol 10	467	0.5	2
Sol 12	500 µL of Sol 11	500	0.25	1

Table S2. Concentration of solutions for the first-series calibration.

Table S3. Concentration of solutions for the second-series calibration

Solution	Volume of solutes	Methanol solvent	Concentration (ng µL ⁻¹)	Mass of standard in 4 µL injected in the tube (ng)
Sol 1			500	2000
Sol 2	100 µL of Sol 1	100	250	1000
Sol 3	100 µL of Sol 2	100	125	500
Sol 4	100 µL of Sol 3	100	62.5	250
Sol 5	100 µL of Sol 4	100	31.25	125
Sol 6	100 µL of Sol 5	100	15.625	62.5
Sol 7	100 µL of Sol 6	100	7.8125	31.25

The settings of the GC–MS for the standard calibrations are shown in Table S4 and were the same for both series.

Sample analysis method	Agilent GC–MSD system (7890A) EI Scan	
Column	Restek 10623 Stabilwax	
Autosampler	Centri	
Injector temperature	250 °C	
Oven temp.	45 °C (5 min), 5 °C min ⁻¹ to 180 °C, 25 °C min ⁻¹ to 250 °C	
Mass range	41–350	
Gas	Constant pressure (working with retention time lock for iso- butylbenzene at retention time of 7.5 min)	
inj	splitless	
Threshold, sampling rate	150, 2	
EMV mode	Relative, 70 eV	
MS source, quad auxiliary temperature	230 °C, 150 °C, 280 °C	
Tube – desorption time desorption temperature	5 min 280 ℃	
Trap – desorption time desorption temperature	3 min (20 °C s ⁻¹ to 300 °C) 300 °C	

Table S4. GC–MS analysis settings for the calibration curve evaluation of selected VOCs.

The calibration curve results are presented in Figs. S1–S5, and are summarized in Table S5.

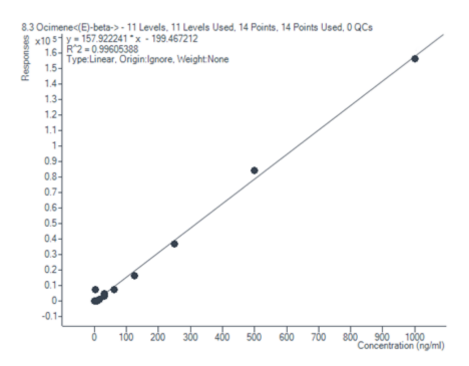


Figure S1. Calibration curve for β -ocimene (E)

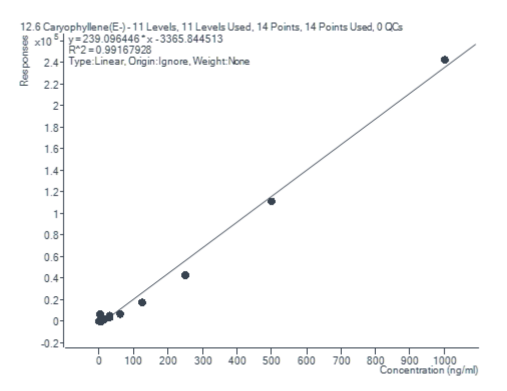


Figure S2. Calibration curve for β -caryophyllene

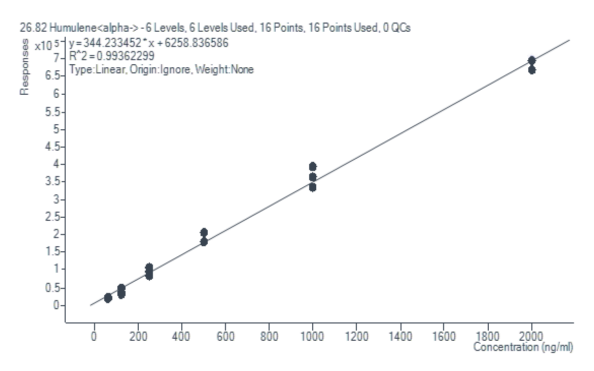


Figure S3. Calibration curve for α-humulene

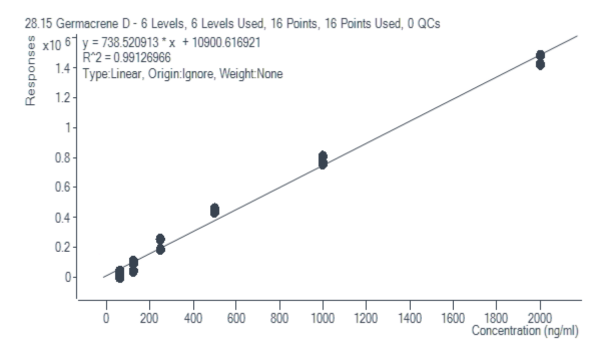


Figure S4. Calibration curve for germacrene D

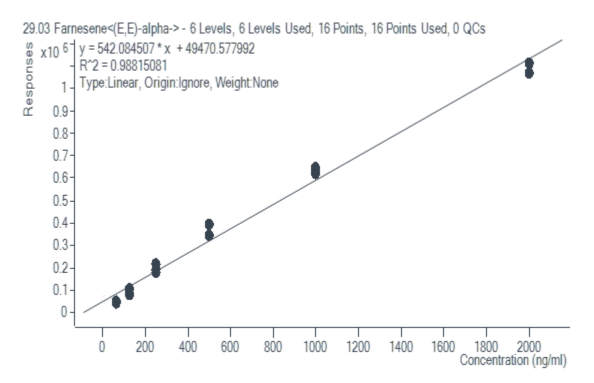
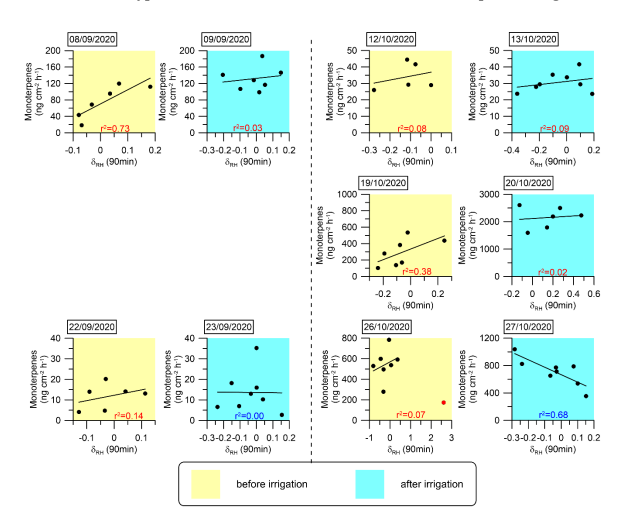


Figure S5. Calibration curve for α -farnesene

Table S5. Regression equa	tions for the five	VOC compounds'	standard calibration curves

VOC compound name	r ²	Regression equation
<i>E</i> -β-ocimene	0.996	y = 157.9220 * x -199.46
β-Caryophyllene	0.991	y =239.096 * x - 3365.84
α-Humulene	0.993	y = 344.233452 * x + 6258.836586
Germacrene D	0.991	y = 738.626951 * x - 4579.005169
α-Farnesene	0.994	y = 617.012200 * x + 22800.770090



S2. Linear and hyperbolic correlation between MTs/SQTs and temporal changes in RH (δ_{RH})

Figure S6. Daily correlations between MT emission fluxes and δ_{RH} . A linear fitting function is used for the fitting curves. The coefficient of determination (r²) for each day is marked in red or blue when the correlation is positive or negative, respectively.

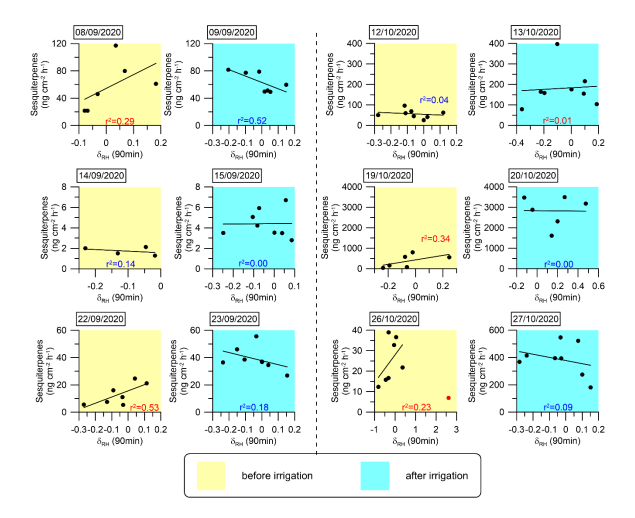


Figure S7. Daily correlations between SQT emission fluxes and δ_{RH} . A linear fitting function is used for the fitting curves. The coefficient of determination (r²) for each day is marked in red or blue when the correlation is positive or negative, respectively. The sample at 12:10 h on 26 Oct 2020 (marked in red) was not considered in the fitting curve for that day, because an extremely sharp increase in RH (from 10 to 31%) occurred within 10 min, which we considered an outlier.

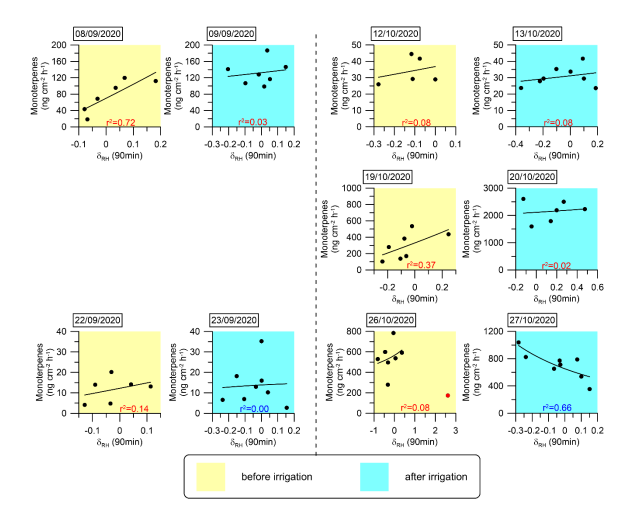


Figure S8. Daily correlations between MT emission fluxes and δ_{RH} . A hyperbolic fitting function was used for the fitting curves. The coefficient of determination (r²) for each day is marked in red or blue when the correlation is positive or negative, respectively.

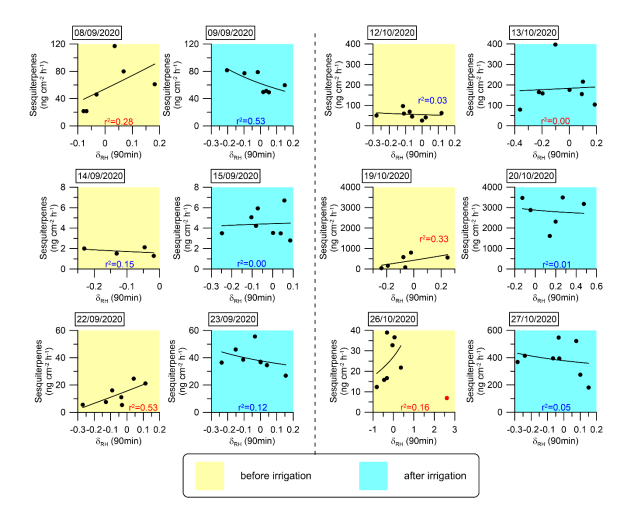
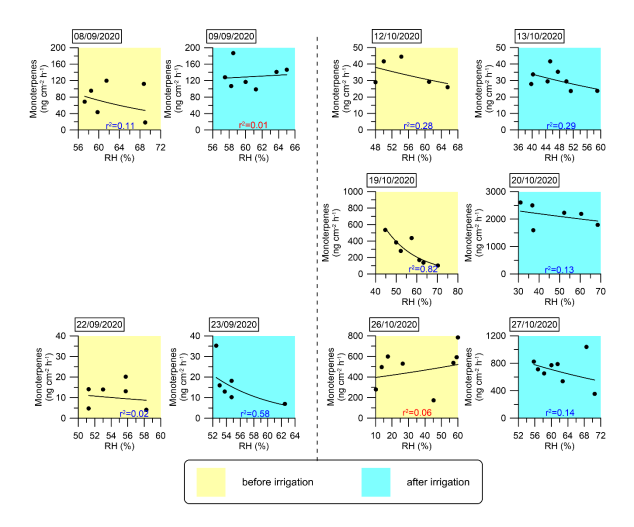


Figure S9 Daily correlations between SQT emission fluxes and δ_{RH} . A hyperbolic fitting function was used for the fitting curves. The coefficient of determination (r²) for each day is marked in red or blue when the correlation is positive or negative, respectively. The sample at 12:10 h on 26 Oct 2020 (marked in red) was not considered in the fitting curve for that day, because an extremely sharp increase in RH (from 10 to 31%) occurred within 10 min, which we considered an outlier.



S3. Exponential correlation between MTs/SQTs and RH

Figure S10. Daily correlations between MT emission fluxes and RH. A linear fitting function was used for the fitting curves. The coefficient of determination (r^2) for each day is marked in red or blue when the correlation is positive or negative, respectively.

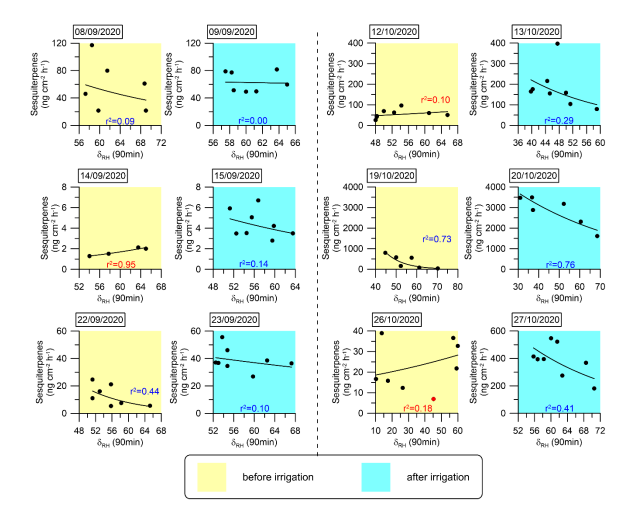


Figure S11. Daily correlations between SQT emission fluxes and RH. A linear fitting function was used for the fitting curves. The coefficient of determination (r^2) for each day is marked in red or blue when the correlation is positive or negative, respectively. The sample at 12:10 h on 26 Oct 2020 (marked in red) was not considered in the fitting curve for that day, because an extremely sharp increase in RH (from 10 to 31%) occurred within 10 min, which we considered an outlier.