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## Supplement of

# Microbial response to deliquescence of nitrate-rich soils in the hyperarid Atacama Desert

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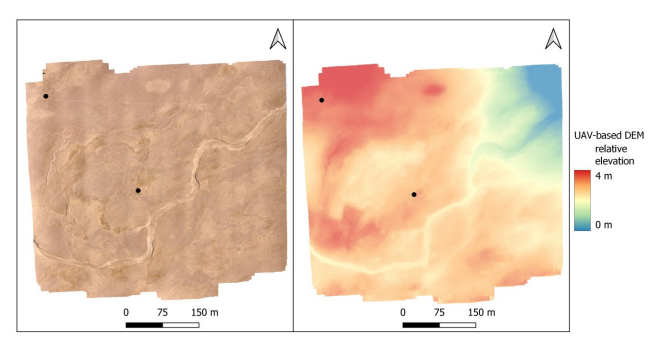


Figure S1: Orthographic image and DEM of the study area of EP1 and EP2

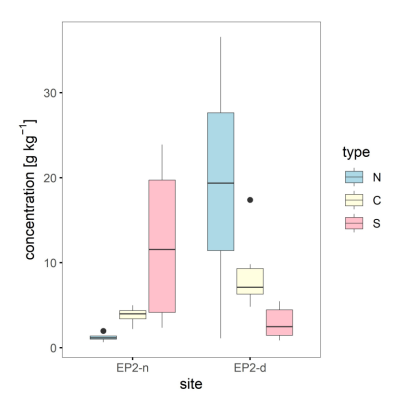


Figure S2: Elemental data (C,N,S) of the EP2 site

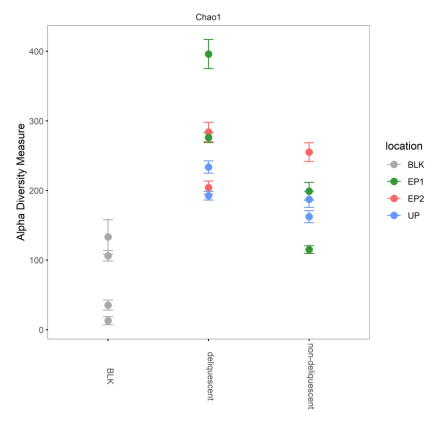


Figure S3: Alpha diversity of the 16S rRNA gene sequencing data

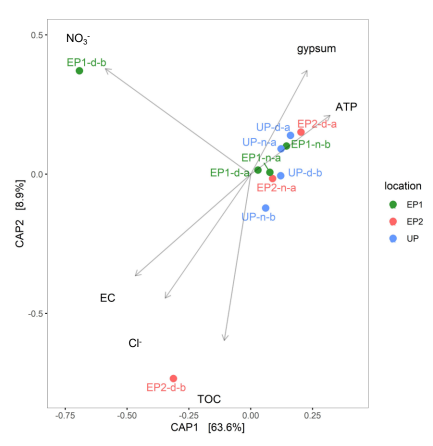


Figure S4: Canonical analysis of Principal Coordinates (CAP) compare the bacterial communities based on 16S rRNA gene sequencing data to different environmental variables (including EC and concentration of gypsum,  $CI^{-}$ ,  $NO_3^{-}$ , ATP, and TOC).

## Compositional profiles of organic molecules

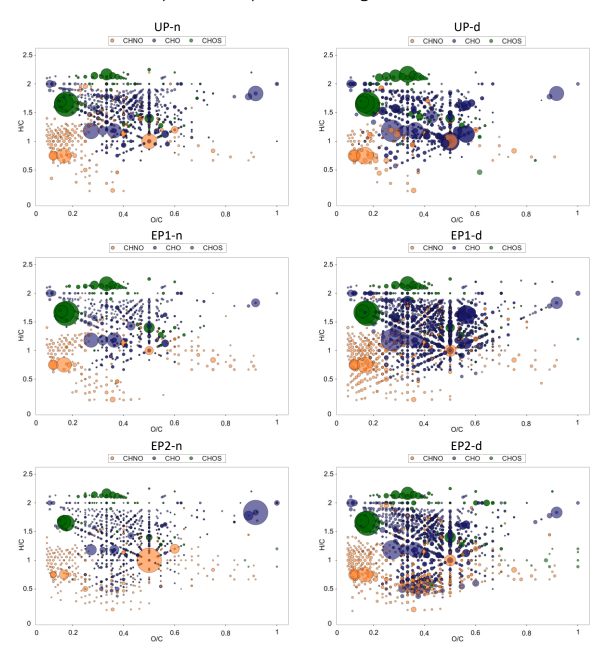


Figure S5: Organic molecule compositional profiles plotted in Van Krevelen diagrams. The hydrogen:carbon atomic ratio (H/C) as a function of the oxygen:carbon (O/C) atomic ratio of organic compounds. The elemental compositions are plotted as CHO (blue), CHOS (green), and CHNO (orange), and bubble sizes depict mass signal intensities.

#### tATP quantification via standard addition method

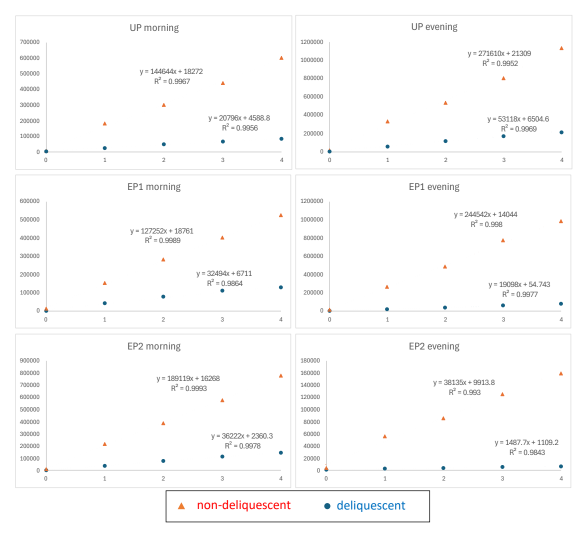


Figure S6: ATP standard addition. Linear fits are plotted for each sample site during morning and evening. y-axis shows the relative light unit (RLU) and the x-axis the standard addition steps in  $\mu$ L. The linear regression function and the coefficient of determination (R²) are plotted above.

#### Calculation:

 $c_x=(b*c_{Sdt})/(m*v_x)$  in mol/L

where  $c_x$  is the ATP concentration,  $v_x$  the sample volume,  $c_{Sdt}$  the concentration of the added standard of the sample. b is the y-intersection and m the slope of the linear regression from the measured standard addition 0 - 4  $\mu$ L 1 $\mu$ M ATP. Linear regression: y=m\*x + b

Table S1 16S rRNA gene sequencing results of the CFUs given in genus level

Phylum	Genus	EP2-d	EP2-n	SUM
Proteobacteria	Pseudomonas	6	0	6
Proteobacteria	Variovorax	0	3	3
Proteobacteria	Agrobacterium	4	5	9
Proteobacteria	Advenella	5	0	5
Proteobacteria	Hydrogenophaga	0	1	1
Bacillota	Bacillus	0	1	1
Actinomycetota	Microbacterium	6	0	6
Proteobacteria	Rhizobium	4	0	4
n sequences		25	10	35