Electronic Supplementary Material (ESI) for Journal of Materials Chemistry B. This journal is © The Royal Society of Chemistry 2017

### **Electronic Supporting Information**

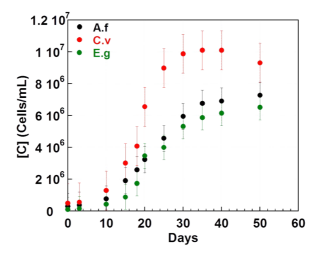
# The physics and chemistry of silica-in-silicates nanocomposite hydrogels and their phycocompatibility

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- **ESI-1**: Experimental details on cell culture
- **ESI-2**: Evolution of gelation time and optical density at 400 nm with total silica concentration at pH 7, 6 and 5.
- **ESI-3**: TEM images of gels at various pH, silicates and Ludox concentrations.
- **ESI-4** Macroscopic observations of cell suspension at various pHs
- ESI-5 TEM images of cells encapsulated at various pH and silicate concentration

#### ESI-1: Experimental details on cell culture

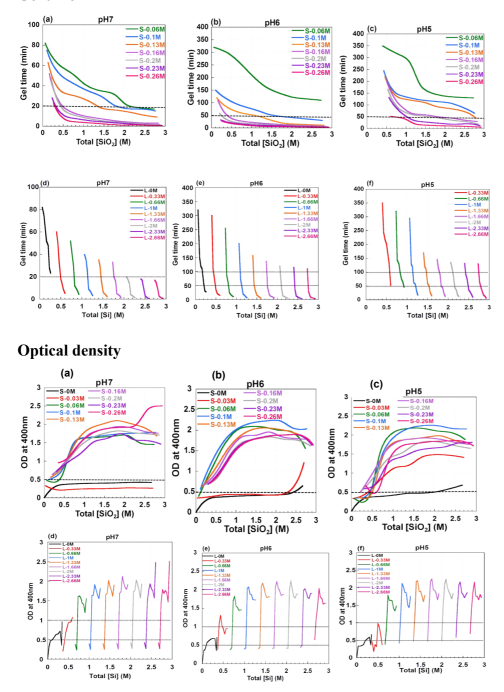
Bold Basal (BB) medium for *A. flos-aquae* and *C. vulgaris* culture and mineral (M) medium for *E. gracilis* culture were prepared according to the literature. All culture media are sterilized by autoclaving (130° C, 20 min, 220 kPa) before use. The micro-algae keeped in erlenmeyer flasks, are placed in growth chamber conditioned to  $(20.0 \pm 1.0)$  °C and is manually shaked once a day at least. The luminosity is adjusted to the optimal intensity for each strain through neon lights (30-60  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> photosynthetic photon flux (PPF) for cyanobacteria, 40-70  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> for green micro-algae). Algae were maintained in nycthemeral cycles of 16 hours of illumination and 8 hours of darkness. Corresponding growth curves are provided below



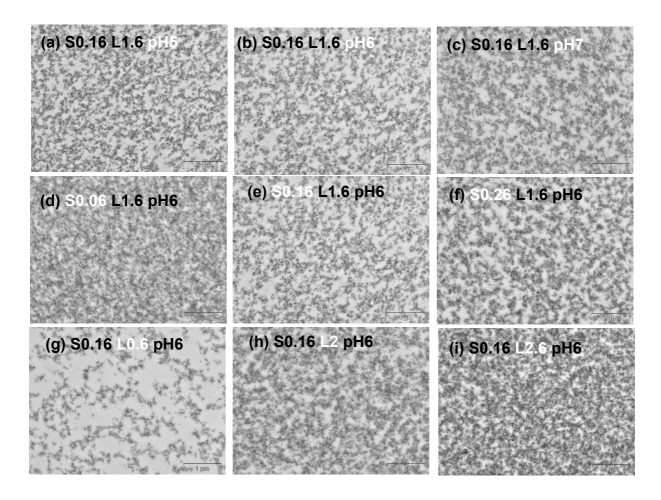
<sup>1</sup>Handbook of Phycological methods. Culture methods and growth measurements. Ed. J. Stein Cambridge University Press. 1973

**ESI-2**: Evolution of gelation time and optical density at 400 nm with total silica concentration at pH 7, 6 and 5: (a-c) each color line is at fixed silicate (S) concentration; (d-f) each color line is at fixed Ludox (L) concentration. Dashed lines represent composition range where similar gel time or optical density can be obtained for different silicate or Ludox concentrations

#### Gel time

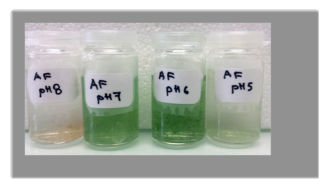


**ESI-3** TEM images of gels (a-c) at fixed silicate and Ludox and variable pH, (d-f) at fixed Ludox and pH and variable silicate, (g-i) at fixed silicate and pH and variable Ludox (scale bar = 1 μm). S = silicate concentration (in M); L = Ludox concentration (in M).



## **ESI-4** Macroscopic observations of cell suspension at various pHs

# Anabaena flos-aquae



Euglena gracilis



Chlorella vulgaris



**ESI-5** TEM images of cells encapsulated in a silica gel at pH 5 ([Ludox] = 1.6 M, [silicate] = 0.16 M) and at [silicate] = 0.26 M (pH 6, [Ludox] = 1.6 M)

