Configurable Compartmentation Enables In vitro Reconstitution of Sustained Synthetic Biology Systems

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Abstract

The compartmentalized and communicative nature of biological cells contributes to the complexity and endurance of living organisms. We present a general compartmentalization strategy for in vitro systems that inherits the passive transport phenomenon of biology. The strategy incorporates layered, micrometer-sized, hydrogel-based compartments featuring configurability in composition, functionality and selective permeability of biomolecules. We demonstrated the utility of the strategy by reconstituting a compartmentalized in vitro protein synthesis system which supports multiple rounds of reactions and compartmentalized living bacteria-based biosensors which sustain long-lasting functioning with markedly enhanced fitness in complex environments. The strategy should be widely applicable for constructing complex, robust and sustained in vitro synthetic molecular and cellular systems, paving the way for their practical applications.
**Introduction**

Compartmentalization is a widespread phenomenon amongst biological systems\(^1\)-\(^4\). The existence of physical barriers within and around cells further enables selective retention of essential biological materials while allowing sustained exchange with outer environments. Through compartmentalization, biological cells possess reduced entropy compared to an open system, which contributes to the high efficiency of cellular machineries. It appears that hierarchical compartmentalization enabled biological systems to gain increasing complexity and diversity during evolution\(^5\). For instance, eukaryotic cells envelop genomic materials and gene transcription activities in the nucleus, which protects the genome while allowing regulation of key cellular events at levels unavailable to prokaryotes\(^6\)-\(^10\). Moreover, many compartments of cells are dynamic and configurable. The permeability of bacteria cells can be tuned to resist environmental impacts through a set of sophisticated mechanisms\(^1\(^{11}\)-\(^1\(^{13}\). The configurable compartmentalization of cells at multiple hierarchical levels contribute to the richness and diversity of biological organisms.

Inspired by nature, we developed a configurable compartmentation strategy for engineered *in vitro* molecular and cellular systems (ConCEiV). Through a microfluidic approach, ConCEiV encapsulates these systems in micrometer-sized, layered hydrogel compartments. Hydrogels are ideal materials for accommodating biomolecules and living cells as they provide a semi-permeable aqueous environment, allowing sustained reagent exchange for diffusion-limited biochemical reactions and cell culture and functioning. The configurability of our hydrogel-based compartments lies in the ease of tuning composition and geometry, of functionalization (e.g. magnetizing) and, importantly, of deterministic adjustment of selective permeability (Fig. 1). These features endow ConCEiV with wide applicability of reconstituting *in vitro* systems across different hierarchies. We demonstrate in this work the compartmentalization of a cell-free protein expression system for sustained, multi-cycle production of target proteins, and of a higher-hierarchy system – an engineered living bacteria-based biosensor exhibiting enhanced environmental fitness.
Over the years, efforts have been devoted to reconstituting cell-like compartments in vitro, with aims to study the complexity of cells from a bottom-up perspective, and to perform highly parallel and heterogenous reactions in a discrete manner. A typical in vitro compartmentalization strategy is water-in-oil (w/o) emulsification which yields isolated aqueous compartments surrounded by an immiscible oil. Coupled in vitro translation and transcription within these w/o droplets has been realized by Tawfik and Griffiths to select desired genes. Although micelle- or vesicle-based transport does allow the exchange of small organic molecules across surfactant-stabilized w/o droplets, many bioactive molecules are hydrophilic and the continuous oil phase barriers molecular transport between compartments and their surroundings. It would be desirable to build a sustained and interactive in vitro system which appreciates an aqueous environment with selectively permeable compartmentalization. Such a system could be living material-based biosensors to monitor environmental signals or bioreactors fueled by external energy/nutrients. In an all-aqueous setting, compartmentalization can be realized by creating membrane-based vesicles such as liposome, proteinosome and polymersome. Some of these membranes are intrinsically porous, or nanopores can be intentionally created to acquire selective permeability. These membrane-based compartments are inherently fragile, limiting their practical adoption in field studies and real-world environments. Moreover, engineering these compartments to meet highly specified and customized permeability requirement remains challenging. With our layered hydrogel-based compartmentalization, ConCEiV provides an elegant solution to these challenges and offers new capabilities that are highly desired in reconstituting in vitro systems.

Synthetic biology is a multidisciplinary area of research that seeks to create new biological parts, devices, and systems. Cell-free protein synthesis (CFPS) is an emerging field in synthetic biology which enables inexpensive and fine control of recombinant protein expression. In CFPS, the physical boundary of cells, cell membrane, is ruptured to form an open in vitro environment, rendering the flexibility for the addition or removal of natural or synthetic components. With ConCEiV, we re-compartmentalize CFPS and artificially tune the selective permeability of compartments to achieve spatially decoupled in vitro transcription and translation (TX-TL): protein-encoding plasmid DNA and RNA
polymerase (RNAP) are confined within the gel membrane, while synthesized messenger RNAs (mRNAs) transport freely to the external buffered solution, where they continue to be translated for producing target proteins. In addition to machineries required for translation, the buffered solution also contains materials and energy for transcription which continuously diffuse into the compartments through the hydrogel membrane. The decoupled TX-TL allows us to preserve transcription within the microgels by recycling plasmids and RNAPs, as well as to replenish translation by feeding fresh solution, thereby enabling multi-cycled, sustained CFPS.

On the other hand, building higher hierarchical systems based on engineered living cells could open up a set of new applications\textsuperscript{27}. The large portfolio of synthetic biology toolkits offers myriads of genetically modified microorganisms (GMMs) with genetic circuits enabling living diagnostics and therapeutics\textsuperscript{31}, programmable living materials\textsuperscript{32}, information storage\textsuperscript{33} and environmental sensing\textsuperscript{34}. However, these functioning genetic circuits exert additional metabolic burden and decrease the GMM’s fitness in a highly complex natural environment\textsuperscript{35, 36}. How may we protect these artificial strains against predators such as bacteriophage, or their fellow microbes\textsuperscript{37}? Furthermore, after they perform each intended task, it would be highly desirable to retrieve these GMMs to extract their recorded information. Currently, while considerable attention has been paid to biocontain GMMs from escaping to natural environments\textsuperscript{38}, protecting GMMs in real-world settings for sustained functioning is being neglected. A competent and deployable protection strategy that improves the fitness of GMMs is yet to be developed. ConCEiV provides a feasible solution to this challenge by compartmentalizing GMMs in layered microgels. The core hydrogel houses the cells and provides a cozy environment for cellular growth and function. The shell hydrogel is configured to have a finer polymer network that barriers external microbial competitors and predators from invading, while simultaneously allows the exchange of nutrition and molecular signals. Bacteria-based living sensors prepared by ConCEiV show near-perfect resistance against wild bacteriophages while maintaining their functionality. We further demonstrate that ConCEiV enables easy retrieval of living sensors after monitoring pollutant levels in synthetic river silt samples and subsequent high-throughput analysis to retrieve recorded pollutant information by flow cytometry.
Results

Manufacturing of ConCEiV compartments

The ConCEiV strategy compartmentalizes in vitro systems within layered microgels through a two-step droplet microfluidic approach (Fig. 2a). First, engineered molecular or cellular species are co-encapsulated with hydrogel solution using a flow-focusing microdevice (Supplementary Fig. 1 a). The generated droplets are then gelled and de-emulsified, yielding monodispersed gel cores. Second, the core beads are reinjected into a dual-junction flow-focusing device (Supplementary Fig. 1b) to build the shell layer. Due to the close-packed nature\textsuperscript{39} of gel beads in microchannels, we are able to collect core-shell hydrogel droplets at ~90% efficiency (Supplementary Fig. 2). Gelling and demulsification are repeated to obtain the final layered hydrogel compartments. The entire procedure can be completed in 3 hours, yielding over 3,000,000 compartments.

Configuring composition and geometry of compartments

The two-step workflow of ConCEiV offers the ease of altering both core and shell hydrogel materials on demand. Fig. 2b shows manufactured core-shell compartments composed of arbitrary combinations of polyacrylamide (PAAm) and agarose hydrogels. PAAm gel has finer polymer network in general, which is favorable for confining biomolecules. Agarose gel features mild radical-free polymerization, thus being more friendly to accommodate living cells. Moreover, the workflow allows precise tuning of the compartment geometry (Fig. 2c). The core diameter and shell thickness are adjustable between 40 – 100 and 5 – 20 μm, respectively, by simple modifications of the microfluidics. If desired, the core of compartments can be selectively dissolved to create an all-liquid cavity suitable for biological reactions. Fig. 2d displays the manufactured compartments featuring a core-shell architecture with each layer labeled by distinct fluorescence. The core gel is reversibly crosslinked which can be completely dissolved within 3 min (Supplementary Fig. 3&4), leading to hollow architecture. The compartments can also be magnetized with ease by involving ferric oxide nanoparticles, thereby enabling convenient recycling and buffer exchange (Supplementary Fig. 5).
Configuring selective permeability of compartments

The selective permeability of hydrogels has been extensively utilized to sieve biomolecules\(^\text{40}\). We hypothesized that the hydrogel shell of ConCEiV compartments could serve as a similar molecular sieve, selectively barring the transport of biomolecules on our demand. To test our hypothesis, we designed a ConCEiV-based polymerase chain reaction (PCR). As illustrated in Fig. 3a, DNA plasmids are pre-encapsulated in the liquid cavities of compartments as PCR templates and assumed to be physically confined therein. A PCR mix is externally introduced and infused into the compartments. If the physical dimension of amplicon DNA (double of its gyration radius \(R_g\)) is greater than the pore size of hydrogel shell \((D_p)\), they are expected to be accumulated in the cavity during the amplification, whereas smaller products are rather free to transport to the outer environment. A preliminary experiment was first conducted, confirming that our customized PCR strategy was able to conduct as designed (Supplementary Fig. 7). We then manufactured a set of compartments having 3 distinct pore sizes of the shell. This was accomplished by tuning the ratio of crosslinker, bis-acrylamide (Bis), of a PAAm gel. Determined by scanning electron microscopy, the average \(D_p\) of a PAAm gel containing 0.6\%, 0.45\% and 0.24\% Bis is 31, 58 and 131 nm, respectively (Supplementary Fig. 6).

We next designed 10 primer pairs targeting amplicons varying from 150 – 1187 bp on the plasmid and performed a series of ConCEiV-based PCRs (Supplementary Fig. 8). Fig. 3b displays the fluorescence distribution of compartments (Supplementary Fig. 9) subject to various PCR conditions and a set of micrographs showing representative 0.6\% Bis-PAAm compartments post PCR. Overall, the DNA amplicons experience a mobility transition from being diffusible through the shell to being confined in the cavity with the increase of their length, which can be noted from the elevated fluorescence profile. For each different \(D_p\), however, the transition occurs at distinct stages; the threshold length at which most of the amplicons can be trapped are estimated to be \(\sim 434, 547\) and \(866\) bp for \(D_p\) of 31, 58 and 131 nm, respectively, in a positive correlation with \(D_p\). These results verify that our compartments indeed possess a size selection effect of DNA molecules, and, importantly, the selective permeability can be readily configured by simple adjustment of the hydrogel shell composition.
It would be beneficial if we could predict the selective permeability of our compartments for any given pore sizes without performing the customized PCRs. With this goal, we built a physical model considering free diffusion of semiflexible DNA molecules in porous media (Supplementary Materials). The model approximates that the diffusible-to-confined transition takes place where the geometrical size of DNA is close to the size of hydrogel pores. In an aqueous solution, the geometry of DNA as a semiflexible polymer can be characterized by its radius of gyration using Kratky-Porod equation\(^1\) (Supplementary Materials, Eq. S1). Based on this, we derived the relation between the critical transition length of DNA and \( D_p \) (Supplementary Materials, Eq. S4). Fig. 3c plots model-defined diffusible and confined regimes divided by Eq. S4. For comparison, we included the experimental data derived from Fig. 3b. As can be seen, the model exhibits reasonable agreement with the experimental results. Therefore, we believe that the established model can provide a preliminary guide of designing selective permeability of our compartments to confine or release desired DNA molecules.

**Establishing ConCEiV-based cell-free protein synthesis**

After examining the configurability of the compartments, we attempted to reconstitute compartmentalized protein synthesis in vitro towards a sustained, long-lasting system. Typical CFPS assays are batch reactions where synthesized protein products and the molecular machineries are all mixed together\(^2\). The latter are not recyclable, often discarded in subsequent protein purification steps, which limits the total cost-effectiveness and disobey with the central goal of synthetic biology for green biosynthesis.

With ConCEiV, we aimed to bridge compartmentalized biological in-vivo protein expression and open-environment CFPS, creating a semi-open protein synthesis system in vitro. As shown in Fig. 4a, ConCEiV-based protein synthesis consists of hollow compartments with their liquid cavities containing genes and enzymes immersed in a feeding solution. We hypothesized that by configuring the selective permeability, protein-encoding plasmids and RNA polymerases (RNAP) could be retained within the compartments, allowing for RNA transcription. Other required materials in the transcription step such as dNTPs and energy can continuously diffuse in from the outer
feeding solution. The synthesized messenger RNA (mRNA) could then transport back to the feeding solution to initiate protein translation.

Having the DNA transport model established (Fig. 3c), we were able to expect that a 2.48 kbp plasmid encoding superfolder green fluorescent protein (sfGFP) would be confined in the compartments. Even taking supercoiling into account does the physical size of the plasmid exceed the largest pore size in our case\textsuperscript{43}. To predict the configurability for protein transport across the compartments, we modified the DNA model by replacing the gyration radius of DNA to that of a globular protein molecule given its total number of amino acids, which yielded a brief model describing the diffusible-to-confined characteristics of proteins (Fig. 4b, Supplementary Materials, Eq. S6). Again, we marked the pore size profile of 0.24\%, 0.45\% and 0.6\% Bis-PAAm hydrogel shell on the plot. The model suggests that T7 RNAP, comprising 883 amino acids, would be confined in 0.6\% Bis-PAAm compartments having an average pore size of 31 nm. By contrast, sfGFP, having only 241 amino acids, could transport through the compartments with the marked pore sizes. It is noteworthy that bacteria ribosome is a huge RNA-protein complex, the number of amino acids of which exceeds 7,000\textsuperscript{44}. Thus, we expect that ribosomes could not transport through all these compartments.

With insights gained from the models, we moved to experimentally constitute ConCEiV-based in vitro protein synthesis systems. We first encapsulated only sfGFP plasmids in the compartments and incubated them in a E. coli extract-based CFPS mixture. The sfGFP fluorescence are weak for all shell pore size conditions (Fig. 4c), suggesting that the TX-TL procedure could barely take place. This is possibly due to the physical barrier effect of the hydrogel shell which limited effective transport of RNAP into the compartment cavity to initiate the transcription. Our model suggests RNAP could transport into the 0.24\% and 0.45\% Bis-PAAm compartments, yet the limited local concentration might lead to extremely low transcription efficiency. Further, we added crude cell extract which contains in-vivo expressed T7 RNAP into the compartment and observed marked increment in sfGFP fluorescence, which suggests that the transcription events indeed occurred in the cavity. Most importantly, the fluorescence level of 0.6\% Bis-PAAm compartments is significantly elevated, 8.5 times of the previous value where only plasmid was added and
over 3.5 times of those compartments with larger pores (0.24% and 0.45% Bis). This reveals a much more efficient transcription occurring within 0.6% Bis-PAAm compartments, suggesting that the 0.6% Bis-PAAm compartments possessed a significantly enhanced confinement effect on RNAPs than other compartments do, in strong agreement with our model prediction. We further derived that the sfGFP yield for 0.6% Bis-PAAm compartments was as high as 0.74 mg/mL, comparable to that of recently reported open CFPS systems\textsuperscript{45}. To further confirm the effect of confined RNAPs on protein expression, we added an extra dose of purified RNAP to 0.6% Bis-PAAm compartments and indeed observed a further increased sfGFP yield (Supplementary Fig. 12). This also suggests that the protein yield of the ConCEiV-based system can be further increased upon additional optimization. These findings verified the successful establishment of our ConCEiV-based in vitro protein synthesis system. As illustrated in Fig. 4d, the system separated transcription and translation in space by confining plasmids and RNAPs in the compartments while allowing the transport of mRNAs to the outer feeding solution, thus achieving a eukaryotic cell-like, spatially decoupled transcription-translation process. To further verify this, we monitored the mRNA level in the feeding solution throughout the protein synthesis procedure (Fig. 4e). The mRNA level of our ConCEiV-based system is comparable to that of bulk CFPS, confirming the reasonably high efficiency of the endo-compartment transcription and efficient transport of the mRNAs to the outer solution.

**Multicycle in vitro protein synthesis enabled by ConCEiV**

Having established an optimal ConCEiV-based protein synthesis system, we proceeded to test whether our proposed goal – sustained and multicycle protein synthesis could be accomplished. In this experiment, we encapsulated sfGFP plasmids and RNAPs in 0.6% Bis-PAAm compartments. The compartments were subsequently immersed in the feeding solution for protein synthesis. After each cycle, the feeding solution was collected and the compartments were magnetically recycled to incubate with new feeding solutions. We repeated 5 cycles and quantified the sfGFP yield for each cycle (Fig. 4f). Although a decay of the expression level was observed along the cycles, possibly due to the deactivation of RNAP after prolonged functioning, a yield of 0.25 mg/mL was still obtained.
at the fifth cycle. The overall yield of the 5-cycled expression exceeded 2.2 mg/mL, around 3 times of its first cycle. These results verified that we have successfully achieved multicycle in vitro protein synthesis, a key step forward towards green and sustainable biosynthesis.

**Establishing ConCEiV-based living biosensors**

We next attempted to compartmentalize a higher-hierarchy system – living bacteria-based biosensors. We envisioned that the ConCEiV system could serve to encapsulate, protect and retrieve GMMs, enabling robust and sustained functioning in complex environments. As a demonstration, we explored the ConCEiV architecture to grow a valerolactam biosensing strain in a core hydrogel matrix, while a tougher semi-permeable outer layer served as a barrier against outside predators, such as phage (Fig. 5a). Lactam is an important class of Nylon precursor and could be detected in the environment as industrial chemical pollutant\(^46\). The concentration of lactam residual in the waste falls within 100 mM range\(^47\). To build lactam-responsive biosensors, we transformed a lactam sensing plasmid into *E. coli* DH5\(\alpha\). The plasmid carries a ChnR-Pb transcription factor-promoter pair that drives mCherry expression upon recognition of lactam by the ChnR protein. We encapsulated the biosensor strain in the 1% agarose core of compartments. 2% agarose gel was used as the shell protective layer. The reconstituted format of lactam biosensors exhibited excellent colony formation and lactam sensing behavior (Fig. 5b). The compartmentalized sensors responded with reasonable linearity between 5 mM and 100 mM valerolactam. These results confirm that the ConCEiV core-shell architecture supports long-term growth and functioning of GMMs.

**Compartmentalized biosensors resisting lytic phages**

We further investigated whether the layered structure of the compartments could provide effective protection against phage attack. Wild DH5\(\alpha\) lytic phages were enriched from swine wastewater samples collected at a farm in Shanghai rural area. A plaque assay was performed, confirming the phage’s lytic activity on the engineered biosensor strain (Supplementary Fig. 14). We then added the phages to the culture medium and/or to the core of the compartments and examined the cell growth. As depicted in Fig. 5c, colonies formed in most of the compartments with the absence of phages. When the phages were
directly added to the core of compartments, the density of colonies dropped significantly, further confirming the phage’s lytic activity. By contrast, the biosensor colonies formed as normal when the phages were only present in the medium. To further confirm the protective effect empowered by the layered ConCEiV architecture, we compared the biosensor growth in bare single-layer (1% agarose, Supplementary Fig. 15) hydrogel beads versus layered ConCEiV system (Fig. 5d). As seen, both systems functioned comparably in the absence of phage. With the presence of phages in medium, however, colonies were found in only less than half of the single-layer beads, whereas for the layered compartments the difference was negligible. Even when phages were present in the core gel over 40% compartments were found positive. By contrast, the positive rate of single-layer beads was less than 10%. This might suggest a bidirectional protective effect of the ConCEiV system that it not only resists the phages from outside but also prevents the leakage of phages from inside, thus limiting the spread rate of phages between the compartments. Overall, these results confirm the excellent resistivity of our ConCEiV-based biosensors against predators from the environment, paving the way of their practical usage in the field.

**Long-lasting functioning and retrieval of biosensors in complex environments**

To further explore the reconstituted biosensors to survive in ill-defined surroundings, to record and memorize desired information, and to be retrieved for reading stored, we designed a polluted river water sensing experiment (Fig. 5e). In the experiment, the biosensor compartments were put in artificial river silt samples polluted by certain levels of valerolactam. After incubation, they were magnetically purified from the samples and then subjected to flow cytometry to extract their stored fluorescence information. Antibiotics experiments reveal the competition between wild microbes and the introduced GMMs (Supplementary Fig. 16). The biosensor growth was found markedly slowed down without the application of strain-specific antibiotics, yet a duration of 32 hours was found sufficient for the biosensors to grow and function. Fig. 5f displays the FACS results of biosensors’ mCherry fluorescence distribution at various pollutant levels. Gaussian fitting was performed to the data, showing that each of the distribution diagrams features 2 distinct peaks. The first peaks fixed at lower fluorescence (5-10 a.u.) were considered
representing empty compartments or debris. Therefore, we discarded them and derived the fluorescence profile from the second peaks as the biosensor output (Fig. 5f, inset). Clearly, the compartmentalized biosensors successfully recorded corresponding valerolactam levels even in such a complex environment. Importantly, the recorded results are consistent with biosensor performance in culture medium (Fig. 5b), suggesting that re-calibration may not be required when the biosensors are adopted to a strange environment. The above results demonstrate a complete round of living cell-based biosensing in a close-to-reality setting.

Discussion

A major advance of ConCEiV is the introduction of configurable semi-permeability to compartmentalized reactors. Realizing such a capability is a daunting task in other established systems such as microdroplets. The hydrogel compartments could directly inherit a vast repertoire of composition-property relationship from decades of hydrogel polymer studies. Applying the classic agarose and polyacrylamide hydrogels, ConCEiV demonstrates compatibility with several in vitro biochemical reaction classes: nucleic acid amplification, TX-TL and cellular growth. The porous barrier mimics the passive transport phenomena in biology and allows replenishable and sustained biological reactions in the parallel compartments. Other functionalization capability of the core-shell system such as magnetization improves handling convenience. Further combination of multiple biochemical steps will allow the reconstitution of even more diverse synthetic molecular and cellular systems. As demonstrated in the present work, the micrometer-sized compartments are readily compatible with commercial analytical tools such as flow cytometers and digital PCR readers. This offers the ease of retrieve stored information of thousands of compartments as independent synthetic biology units.

While our strategy possesses notable advantages as demonstrated above, the current manufacturing scheme might impose several limitations. First, the two-step encapsulation could result in imperfect compartment formation. Although the internal cores can be coated by hydrogel shells with established “beating-Poisson” microfluidic techniques, such a procedure can hardly reach 100% efficiency. Second, the compartment could occasionally break, therefore releasing biological materials in the cavity to the outer
environment, ending up with competition for reaction substrates. Third, unlike liposome or cellular membrane, where the core and the outer environment could maintain different liquid composition, the permeable nature of the compartment precluded some biological processes involving active transport. Besides, the tuning of selective permeability solely relies on adjustment of hydrogel composition in the current format. It would be favorable if the permeability could be controlled by more active means such as temperature or light. This demand can be potentially met by incorporating temperature- or photo-responsive hydrogel materials.

Owing to its highly configurable nature, the ConCEiV system holds great potential for future synthetic biology applications. For instance, it is possible to encapsulate a large diversity of individual mutants, while performing dedicated reactions separately for directed evolution. As a physical barrier, the compartment could offer protection for the encapsulated living organisms in native environment, such as human gut, while allowing nutrient exchange with the surroundings. The development of such tools could be helpful to in-vivo microbiome studies of unculturable bacteria and living bacteria-based therapeutics.


Figure 1. Schematic of the ConCEiV strategy. ConCEiV compartmentalizes open in vitro molecular and cell-based systems in micrometer-sized, layered hydrogel-based compartments. The compartments own configurability in selective permeability, composition, geometry and functionalization. The compartmentalized systems allow biological reactions and activities executing in a sustained manner due to configured passive transport. For a reconstituted molecular system, the selective permeability of compartments can be tuned to achieve desired transport of biomolecules such as protein and DNA while allowing constant input of reactants and energy. A reconstituted cell-based systems can protect the encapsulated engineered cells from competitors and predators and meanwhile support the growth and functioning of the cells, thereby enhancing the fitness of these functional synthetic biology systems in complex environments.
Figure 2. Manufacturing of layered hydrogel-based compartments. (a) Microfluidic workflow to encapsulate engineered synthetic biology systems in layered compartments. The systems are pre-encapsulated in the hydrogel cores using a microfluidic dropmaker. The core beads are further reinjected to a second microfluidic device to generate layered compartments. (b-d) Optical micrographs of (b) compartments composed of arbitrary combinations of polyacrylamide (PAAm) and agarose hydrogels, (c) compartments featuring tunable core diameters between 40 to 100 μm and shell thickness between 5 to 20 μm and (d) core-shell hydrogel compartments and hollow compartments after core dissolving. The core and shell layer of compartments in (d) are fluorescently labeled with Cy3 (red) and FITC-dextran (green), respectively. Scale bars: 50 μm.
Figure 3. Configuring selective permeability of ConCEiV compartments. (a) Schematic of a compartmentalized PCR assay designed to explore the selective permeability of compartments. If the dimension of amplicon DNA (double of its gyration radius $R_g$) is greater than the pore size of compartment shell ($D_p$), they are expected to be accumulated in the cavity. By contrast, smaller amplicons will diffuse to the outer solution. The process can be visualized by fluorescent labeling of the amplicons. (b) Results of compartmentalized PCR performed in compartments with configured pore sizes. The scatter plots (top) display fluorescence profile (normalized) of the compartments subject to serial PCRs. Each point in the plots represents the value of a single compartment. The series of fluorescent micrographs (bottom) show individual 0.6% Bis-PAAm compartments undergone corresponding PCR conditions. Scale bar: 50 μm. (c) Physical model describing the transport characteristic of dsDNA across the compartments. The scattered points represent experimental results (legend) obtained from the compartmentalized PCR. The shadowed areas represent the extent of standard deviation of $D_p$ calculated from SEM measurements (Supplementary Fig. 6).
Figure 4. ConCEIV-based in vitro protein synthesis. (a) Schematic of the compartmentalized protein synthesis system. Hollow Bis-PAAm compartments encapsulating plasmid and RNA polymerase (RNAP) are immersed in a feeding solution containing other required reagents for RNA transcription and protein translation. (b) Physical model describing the transport characteristic of globular proteins across the compartments. The dashdotted lines represent the averaged pore size of three different Bis-PAAm gels with corresponding shadowed areas denoting the extent of the standard deviation. Vertical dotted lines indicate total amino acid number of the stated protein molecules. (c) sfGFP fluorescence levels (normalized) of protein synthesis assays performed on three different Bis-PAAm compartments. In the 1st group (bright background) of experiments, solely sfGFP plasmid was added to the compartment cavity. *E. coli*
BL21 cell extract containing *in-vivo* expressed T7 RNAP was added along with sfGFP plasmid in the 2nd group (grey background). Error bars denote standard deviations of multiple parallel experiments (n ≥ 8). Statistical analysis: unpaired two-tailed t-test. (d) Proposed transcription-translation procedure in the ConCEiV-based system. Transcription and translation are spatially separated. Plasmids and RNAPs are confined in the compartments for transcription. The produced mRNAs transport to the feeding solution where translation takes place. (e) Total mRNA levels during sfGFP expression by a ConCEiV-based system and a bulk CFPS system. Error bars denote standard deviations of multiple parallel experiments (n = 3). (f) Workflow (left) and yield (right) of multicycle sfGFP synthesis by a ConCEiV-based system. For each cycle the 0.6% Bis-PAAm compartments containing sfGFP plasmid and cell extract were incubated in a fresh feeding solution for 6 hours, after which compartments were magnetically recycled and the sfGFP yield in the supernatant was quantified.
Figure 5. ConCEIV-based living biosensors. (a) Schematic of the compartmentalized biosensor. An engineered E. coli strain is encapsulated in the cores (1% agarose) of layered compartments for cell growth and functioning. The strain senses an industrial chemical pollutant – lactam through a constructed lactam-mCherry pathway. A dense outer layer (2% agarose) protects the encapsulated biosensor strain from environmental impacts such as phage attack. (b) Characterization of the compartmentalized biosensors. Top: fluorescence of individual biosensor compartments (n ≥ 50) versus incubation time. The compartments were cultured in medium containing 100 mM valerolactam. Bottom: dose-response relation of the compartmentalized biosensors (n ≥ 50). (c) Optical micrographs showing growth condition of biosensor strain encapsulated in core-shell compartments in presence of phage. The compartments were incubated in medium containing 5 mM valerolactam for 16 hours. The bacteria colonies are labeled in red as they express mCherry fluorescent protein. (d) Comparison of biosensor strain growth in single-layer agarose (1%) beads and in double-layer compartments in presence of phage. Microgels having grown colonies were considered positive. Statistical analysis: unpaired two-tailed t-test. (e) Workflow of sensing, retrieval and analysis of the compartmentalized biosensors in an artificial river silt environment polluted by lactam. (f) Fluorescence profile of compartments after sensing varying levels of valerolactam in polluted river silt. Dashed curves denote dual-peak Gaussian fitting. The micrographs display representative compartments at relevant biosensing conditions. The inset scatter plot shows the mean and standard error of corresponding sensing results. Scale bars: 100 μm.
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Competing Interests: The authors have filed a provisional patent based on this work.

Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper are available from authors upon reasonable request.