Translation-coupled RNA replication and parasitic replicators in membrane-free compartments†

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We report RNA self-replication through the translation of its encoded protein within membrane-free compartments generated by liquid–liquid phase separation. The aqueous droplets support RNA self-replication by concentrating a genomic RNA and translation proteins, facilitating the uptake of small substrates, and preventing the replication of parasitic RNAs through compartmentalization.

Bottom-up construction of an artificial cell is a viable approach to dissect the roles and mechanisms of cellular functions and provide novel cell-like reactors that can be readily manipulated.1–5 To better mimic cell behaviour and create complex biological systems, a key step in artificial cell construction is the compartmentalization of gene expression machinery, a central part of living cells that converts genetic information into proteins to facilitate cellular processes. Typically performed in artificial cells with phospholipid membranes, previous research integrated cell-free gene expression systems with diverse cellular phenomena including quorum sensing,6 cell deformation,7,8 energy generation,9 and replication of DNA and RNA.10,11 A phospholipid membrane, however, strictly limits the transport of molecules to and from compartments, which restricts the applicability of artificial cells. An alternative way to create artificial cells is to use membrane-free compartments, such as cell-like hydrogels and droplets based on liquid–liquid phase separation (LLPS). Although the expression of reporter genes was achieved in these systems,12–17 the demonstration of complex translation-activated biological processes has been challenging. Because membrane-free compartments are generally highly permeable, designable to sequester specific biomolecules, and molecularly crowded like living cells, they could provide unique regulation towards gene-expression and integrated biological reactions.3,5

LLPS provides membrane-less cell-like structures that are ubiquitously seen in natural cells as biomolecular condensates.18 The liquid-like state enables molecular exchange with environments and the selective sequestration of biomolecules, while maintaining the mobility of sequestered biomolecules.19,20 The development of such cell models could help to elucidate the roles of cellular condensates as well as to establish dynamic artificial cells. Because LLPS occurs by simple physical processes, it would also provide insights into primitive compartmentalization by phase separation at the origins of life, especially before the advent of membranes.21,22 To date, enzymatic and catalytic reactions, including the expression of fluorescent reporters, were demonstrated in two types of LLPS-based protocell models, coacervates12,15,23–25 and aqueous two-phase systems (ATPS).13,26–28 The integration of gene-expression with downstream biological processes in these systems, if achieved, would expand the versatility of LLPS-based droplets as artificial cells.

Here, we employ cell-free translation within an ATPS and demonstrate the self-replication of a genomic RNA by its encoded protein. The replication of genetic molecules using self-encoded proteins is essential in cells, allowing autonomous proliferation as well as Darwinian evolution.29,30 We further show additional roles of ATPS in facilitating the sustained replication of the genomic RNA: the prevention of parasitic RNA replication and the facile uptake of building blocks.

To demonstrate the translation-coupled RNA replication (TcRR) in ATPS, we combined a TcRR system based on previous studies29,31 with an ATPS of polyethylene glycol (PEG, 20 kDa, 15 wt%) and dextran (DEX, 9–11 kDa, 1.5 wt%) (Fig. 1A). The TcRR system consists of an artificial single-stranded genomic RNA (2041 nt) that encodes the core subunit of Qb replicase and a reconstituted E.coli translation system based on the PURE
In 8 mM Mg(OAc)$_2$, the genomic RNA replicated 4-fold in (A) 16 mM or (B) 8 mM Mg(OAc)$_2$ in the presence or absence of the ATPS, or in a continuous 1.5 wt% DEX 9–11 kDa or 15 wt% PEG 20 kDa phase. The RNA concentration was measured by quantitative RT-PCR. The dashed lines represent 1-fold replication (RNA concentration after the reaction divided by initial RNA concentration). (c) The TcRR reaction with diluted translation proteins (Table S1, ESI†) in the absence or presence of the ATPS (16 or 8 mM Mg(OAc)$_2$, respectively). All error bars indicate standard errors (N = 3).

Next, we performed the TcRR reaction at 37 °C for 2 h in various conditions and determined the respective level of RNA replication. In the standard TcRR condition optimized without an ATPS (Table S2 ESI† except with 16 mM magnesium acetate (Mg(OAc)$_2$)), the replication of the genomic RNA was detectable only in the absence of the ATPS (Fig. 2A). We optimized the reaction condition with the ATPS and found that changing the concentration of Mg(OAc)$_2$ from 16 mM to 8 mM significantly improved the TcRR reaction in the ATPS (Fig. S1, ESI†); similar DEX-rich phase droplets appeared to form in both Mg(OAc)$_2$ conditions (8 mM in Fig. 1 and 16 mM in Fig. S2, ESI†). In 8 mM Mg(OAc)$_2$, the genomic RNA replicated 4-fold in the ATPS, whereas the replication was negligible in the absence of one or both of DEX or PEG (Fig. 2B). We noticed that the TcRR reaction was detectable only if we mixed the translation system with DEX before the addition of PEG, perhaps because DEX keeps translation proteins soluble and functional despite the high concentration of PEG; PEG is known to inhibit cell-free translation probably due to protein precipitation. The size of DEX-rich phase droplets remained in a similar range after the incubation, although the number of smaller droplets diminished (Fig. S3, ESI†). Furthermore, in contrast to the bulk-optimized condition, the concentration of translation proteins in the DEX-rich phase droplets of the ATPS (Fig. 1C) allowed the dilution of the set of translation proteins (Table S1, ESI†) up to 4-fold without diminishing RNA replication (Fig. 2C).

One of the roles of compartments in prebiotic evolution is to allow sustainable replication of genetic replicators by limiting the propagation of parasitic ones. To examine whether the ATPS could shelter the genomic RNA from parasitic RNAs despite the lack of membranes, we prepared two types of droplets, which contained either the genomic RNA or a 510 nt parasitic RNA (Parasite-γ115†). The parasitic RNA lacks the replicase subunit gene, but it can replicate by exploiting the replicase expressed from the genomic RNA (Fig. 3A). We first encapsulated the Cy5-labeled genomic RNA and the Cy3-labeled parasitic RNA in distinct ATPS droplets (Fig. S4A, ESI†). By gently mixing the two ATPS, we examined the inter-droplet diffusion of the two RNAs. Although some droplets showed both fluorescence signals, the distribution of the fluorescence-labelled RNAs was not uniform across the droplet population, and one type of RNA dominated in most of the droplets (Fig. 3B). The uneven RNA distribution among the droplets...
**Fig. 3** The TcRR reaction in the presence of parasitic RNAs. (A) The parasitic RNA replicates by utilizing Qβ replicase expressed from the genomic RNA. (B) Confocal microscope images immediately after mixing two separately prepared ATPS, containing Cy5-labelled genomic RNA or Cy3-labelled parasitic RNA. The ratio of the two ATPS was 1:1 for visualization. Left panel: Cy5 fluorescence. Middle panel: Cy3 fluorescence. Right panel: Both fluorescence channels merged with the transmitted light image. (C) A merged image after 2 h incubation at 37 °C. All scale bars = 30 μm. (D) Two ATPS, containing one, both, or none of the genomic and parasitic RNAs, were mixed as illustrated in a 19 : 1 (a) : 1 (b) ratio (#1 – #3). The ATPS and bulk (#4) mixtures were incubated at 37 °C for 2 h, followed by quantitative RT-PCR to measure RNA concentrations. Initial concentrations of the genomic and parasitic RNAs were 3 nM in all cases. (E) 10 nM of a genomic RNA was incubated at 37 °C for 2 h in the ATPS (#5) or bulk (#6), followed by agarose gel electrophoresis. The concentration of minus-strand genomic RNA was measured by quantitative RT-PCR. For all reactions in bulk or the ATPS, 8 or 16 mM of Mg(OAc)₂ was used, respectively. All error bars show standard errors (N = 3–5).

We then performed the TcRR reaction by encapsulating the genomic and parasitic RNAs together in the same ATPS or by gently mixing two ATPS, each containing one type of RNA (Fig. 3D). When we pre-mixed the genomic and parasitic RNAs in the same ATPS (#1), the genomic RNA barely replicated due to the high replication of the parasitic RNA. However, when mixing two ATPS that separately contained the genomic and parasitic RNAs in a 19:1 ratio (which equalized the concentration of the two RNAs in the mixture) (#2), the genomic RNA replicated at the same level as the control reaction performed with an ATPS in the absence of the parasitic RNA (#3), whereas the replication of the parasitic RNA was inhibited compared to the reaction performed with the pre-mixed genomic and parasitic RNAs (#1). We also performed the reaction with a mixed genomic and parasitic RNA system in the absence of ATPS (#4), and found that the replication of the genomic RNA was more efficient with the ATPS droplets, with a significant suppression of replication of the parasitic RNA (#2). These results suggested that the ATPS could support the RNA self-replication by compartmentalizing the genomic and parasitic RNAs in distinct droplets.

In the evolution of genomic RNAs, parasitic RNAs likely appear through mutations in compartments where a genomic RNA already exists.36,37 We therefore also examined whether the ATPS could prevent the replication of a small amount of parasitic RNA (de novo synthesized or contaminated) without manual segregation as performed in Fig. 3D. As described in the Supplementary Results section (ESI†), we found that the ATPS prevented such parasitic RNA (∼220 nt) replication (Fig. 5S, ESI†). Moreover, in the TcRR reaction without the addition of known parasitic RNAs, a genomic RNA (R3019) replicated slightly better in the ATPS (#5, Fig. 3E) than in bulk (#6) by preventing the appearance of parasitic RNAs, even with the inhibitory ATPS condition (Fig. 2A and B). These results emphasize the effectiveness of the membrane-free compartments to support RNA self-replication by limiting parasitic RNA replication.

While the ATPS appeared to prevent inter-droplet RNA diffusion (Fig. 3), LLPS-based droplets are generally permeable to small molecules.19,20 We therefore wondered whether the TcRR reaction could continue by externally supplying small substrates. We initiated the TcRR reaction in the ATPS with 62.5 μM cytidine triphosphate (CTP, used in RNA polymerization), and then added 625 μM CTP to the ATPS after 2 h incubation and continued the reaction. The extra CTP added was as low as 0.03 volume of the ATPS so that the composition of the ATPS was basically unchanged. While in the absence of additional CTP a genomic RNA replicated only up to the resource limitation (corresponding to approximately 120 nM genomic RNA), the TcRR reaction with added CTP exceeded this limitation and used up nearly the entire of the added CTP by 5 h (Fig. 4), suggesting that the ATPS droplets were fully permeable to substrates, with the CTP being supplied to the droplets in a dosedependent manner.
protein (Fig. 2B). Two characteristics of LLPS-based droplets, \(^{19,20}\) concentration of internalized components and permeability to small molecule analytes, allowed us to reduce translation proteins without hindering the TeRR reaction (Fig. 2C) and to supply substrates externally to continue the reaction (Fig. 4). Such a persistent genome replication system functional with a lower input of translation proteins would be a step toward realizing a self-regenerating and long-lasting artificial cell.\(^4\)

Moreover, the ATPS supported the RNA self-replication by sheltering a genomic RNA from various short parasitic RNAs (Fig. 3 and Fig. S5, ESI\(^1\)). The primary mechanism for this effect was to segregate the parasitic RNAs into distinct droplets and prevent inter-droplet RNA diffusion. Such effective compartmentalization is a key feature of cell-like structures for the evolution of genetic entities and previously shown only in compartments with explicit boundaries.\(^{29,30,36,37}\) Therefore, future studies should examine whether the functional RNA is evolvable even in membrane-free compartments, an attractive question in the field of origins-of-life. More discussions can be found in the Supplementary Discussion (ESI\(^1\))

In conclusion, we demonstrated translation-integrated RNA replication within DEX/PEG ATPS droplets, where the replication machinery was concentrated, accessible to external building blocks, and protected from takeover by parasitic RNAs. These results validate LLPS-based droplets as artificial cells for implementing complex biological reactions as well as model systems to study a primitive type of compartmentalization.

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Conflicts of interest

There are no conflicts to declare.

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