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A synthetic replicator drives a propagating reaction-diffusion front

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ABSTRACT: A simple synthetic autocatalytic replicator is capable of establishing and driving the propagation of a reaction-diffusion front within a 50 µL syringe. This replicator templates its own synthesis through a 1,3-dipolar cycloaddition reaction between a nitrone component, equipped with a 9-ethynylanthracene optical tag, and a maleimide. Kinetic studies using NMR and UV-Vis spectroscopies confirm that the replicator forms efficiently and with high diastereoselectivity and this replication process brings about a dramatic change in optical properties of the sample – a change in the color of the fluorescence in the sample from yellow to blue. The addition of a small amount of the pre-formed replicator at a specific location within a microsyringe, filled with the reaction building blocks, results in the initiation and propagation of a reaction-diffusion front. The realization of a replicator capable of initiating a reaction-diffusion front provides a platform for the examination of interconnected replicating networks under out-of-equilibrium conditions involving diffusion processes.

The spontaneous generation of stationary patterns and propagating fronts in chemical systems has intrigued scientists for generation. Such phenomena are ubiquitous in nature and the physical processes behind their appearance and stability have been studied extensively and are now relatively well understood. Propagating reaction-diffusion fronts have received significant attention in this respect. Frequently, one or more oscillatory or autocatalytic processes are found at the core of these systems. Front generation is initiated when an autocatalyst is added at a discrete location in an expansive reaction, initially at uniform concentration and the ensuing reaction generates wave fronts, which propagate outward from the initial reaction zone. In almost all of the examples reported to date, the autocatalysis is based on inorganic chemistry, although, more recently, a small number of examples based on RNA and DNA have been described. Self-replication represents a niche of autocatalytic behavior in which a structurally complex template is capable of recognizing the building blocks necessary for its own formation and catalyzing their reaction to form an exact copy of itself. We, and others, have described the use of such systems in instructable networks, as tools for dynamic systemic resolution and in the construction of mechanically-interlocked molecules. Although all of these systems display the nonlinear kinetic characteristics of autocatalytic systems to a greater or lesser extent, in general, they have been studied under well-stirred batch reactor conditions. The consequence of this reaction format places a fundamental limit on the level of complexity and emergence that can be generated by such system. In order to create diverse emergent behavior there is a need to study self-replicating systems under out-of-equilibrium conditions and propagating reaction-diffusion fronts could provide an ideal vehicle for such studies. Here, we report the design and implementation of a molecular replicating system capable of generating and sustaining a propagating reaction-diffusion front.

Previously, we have described an efficient synthetic replicator based on the general design shown in Figure 1a. Reaction of nitrone 1a with maleimide 2 in CDCl3 at −10 °C results in the rapid, autocatalytic formation of the cycloadduct 3a. Cycloadduct 3a is a very efficient template for its own formation – it is capable of accelerating the reaction between 1a and 2 up to 125 × through a ternary complex [1a•2•3a] and the structure of this complex ensures that only the trans diastereoisomer of 3a is formed during this process. Conventionally, we have monitored the kinetics of replication processes by NMR spectroscopy. In a reaction diffusion format, this reaction requires an alternative method to monitor the progress of the reaction. Ideally, we desired an optical signature of replication. Therefore, we designed nitrone 1b, bearing a 9-ethynylanthracene tag. RM1 calculations (Figure 1b) indicated that this nitrone, in partnership with maleimide 2, was capable of furnishing template 3b through a ternary complex [1b•2•3b]. This complex should permit the formation of only the trans diastereoisomer of 3b. TD-DFT calculations (see Supporting Information) indicated that a significant change in the 350 to 400 nm region of the UV-Vis spectrum could be expected on conversion of nitrone 1b to cycloadduct 3b.
as a result of the presence of the 9-ethynylantracene unit. We synthesized nitrone 1b using standard methods and this compound forms yellow-colored solutions in CDCl$_3$, which exhibit an intense yellow fluorescence (Figure 1a). Pleasingly, the conversion of 1b into 3b resulted in a very significant color change – the yellow fluorescence of 1b being replaced (Figure 1a) by the blue fluorescence of 3b.

![Figure 1](attachment:image1.png)

Figure 1. (a) A self-replicating template 3a is constructed by the reaction of nitrone 1a with maleimide 2 through in the catalytically-active [1a•2•3a] ternary complex. Replacement of the substituent R affords replicator 3b, which incorporates a 9-ethynylantracene optical tag, derived from nitrone 1b. The formation of cycloadduct 3b, mediated by ternary complex [1b•2•3b], is now associated with a change in fluorescence from yellow (1b) to blue (3b). (b) Calculated (RM1) structure of the transition state leading to 3b from the ternary complex [1b•2•3b].

Next, we conducted a series of kinetic experiments involving the reaction of 1b and 2 in CDCl$_3$ at 0 and 20 °C, monitoring the production of cycloadduct 3b by 500 MHz $^1$H NMR spectroscopy. The results of these experiments and subsequent fitting of the experimental data at 20 °C to the appropriate kinetic model are shown in Figures 2a and 2b. These kinetic experiments reveal that replicator 3b is an excellent template for its own formation – the ternary complex [1b•2•3b] generates an effective molarity$^{13}$ (EM) of 16.2 M for the cycloaddition reaction (for details, see Supporting Information). This value for the ternary complex EM is broadly similar to those determined previously$^{9e,10b,11}$ for similar replicators and indicates that the incorporation of the 9-ethynylantracene optical probe has essentially no effect on the functioning of the replicator.

![Figure 2](attachment:image2.png)

Figure 2. (a) Concentration (red circles) and rate (black dotted line) vs time profile for the formation of 3b from nitrone 1b and maleimide 2 as determined by 500 MHz $^1$H NMR spectroscopy ([1b] = [2] = 10 mM, 20 °C, CDCl$_3$). The red line shows the fit of the appropriate kinetic model to these data. (See Supporting Information) (b) The appearance of the resonance associated with the formation of the trans-3b cycloadduct over time in the 500 MHz $^1$H NMR spectrum of a reaction mixture containing 1b and 2 ([1b] = [2] = 10 mM, 20 °C, CDCl$_3$). (c) Selected UV-Vis spectra recorded during the reaction of 1b and 2 ([1b] = [2] = 10 mM, 20 °C, CDCl$_3$), showing the disappearance of nitrone 1b (346 nm band) and simultaneous appearance of cycloadduct 3b (297 nm band). (d) Comparison of absorbance at 297 nm vs time. The blue circles show data determined experimentally from the spectra in Figure 2c. The red line shows the absorbance at 297 nm vs time computed using the concentrations of 3b determined from the best fit of the appropriate kinetic model to the NMR data in Figure 2a.

Having established that 3b was indeed capable of templating its own formation, we next sought to establish that the color change that is observed during this reaction is a signature of the autocatalytic replication processes. Accordingly, we monitored the formation of 3b using UV-Vis spectroscopy (Figure 2c) under identical conditions to those employed in the NMR kinetic exper-
iments. As expected, the UV-Vis spectra recorded during the reaction show the disappearance of a band corresponding to nitrone 1b at 346 nm with the concomitant appearance of a band at 297 nm corresponding to cycloadduct 3. In order to relate this data to the kinetic data derived from NMR spectroscopy, we reconstructed the reaction profile at 297 nm by computing the expected absorbance at this wavelength from the concentrations of the components of the reaction mixture determined from the best fit to our kinetic model to the NMR data. The excellent agreement (Figure 2d) between the calculated and observed reaction profiles provides compelling evidence that the color change that is observed during this reaction is indeed the signature of the replication of 3b.

Figure 3. (a) Graphical representation and (b) photograph of the experimental setup employed for investigation of the propagating reaction-diffusion front initiated by replicator 3b in two 50 µL gas tight syringes. The upper syringe in each case represents the control experiment comprising the nitrone 1b and maleimide 2 only. ([1b] = [2] = 5 mM, 20 °C, CDCl3). The lower syringe is seeded with ca. 2 µL of a solution of 3b. ([1b] = [2] = 5 mM, [3b] = 10 mM, 20 °C, CDCl3).

Normally, propagating reaction-diffusion fronts are observed in reactions that are initiated on flat plates or within capillary tubes. Since CDCl3 is a relatively volatile solvent, we chose to investigate whether 3b was capable of supporting a propagating reaction-diffusion front within a 50 µL gas tight syringe of internal diameter 1.03 mm. Figure 3 illustrates our experimental setup. Two syringes were placed side-by-side in a specially constructed stand housed within a controlled environment where the temperature was regulated at 20 °C. One syringe was filled with a 5 mM solution of nitrone 1b and maleimide 2 in CDCl3. The second syringe was prepared identically with the exception that approximately 2 µL of a 10 mM solution of replicator 3b was drawn into the end of the syringe after it was filled with the solution of 1b and 2.

We envisaged that the syringe containing only 1b and 2 would change color uniformly as replicator 3b was formed. In the other syringe, the presence of 3b would initiate the replication process and the diffusion of the replicator thus formed would establish a reaction-diffusion front that would propagate along the syringe, being observed as the progression of a blue band along the initially yellow syringe.

Figure 4. (a) Processed grayscale images, acquired over time with the syringes illuminated using a 365 nm UV lamp, of the template-initiated reaction-diffusion experiment (+3b, left column) and the control experiment (–3b, right column). (b) Smoothed profiles of the grayscale images from the reaction-diffusion experiment (+3b) over time showing the progression of the reaction diffusion front over 20 minutes.

The syringes were illuminated using a 365 nm UV lamp and a color image was captured every two minutes using a digital camera. Processing of these images (see Supporting Information) afforded the data shown in Figure 4a. It is evident from these images that replicator 3b has, established (Figure 4a, +3b) a propagating reaction-diffusion front within the syringe to which it was added initially. By contrast, no such feature is evident within the control syringe (Figure 4a, –3b). As an additional control, we examined the diffusion of 3b in the absence of an autocatalytic reaction. Approximately 2 µL of a 10 mM solution of replicator 3b was drawn into the end of a syringe filled with a 5 mM solution of nitrone 1b only in CDCl3. In this case, the blue color of replicator
3b disappears as a result of diffusion processes within the syringe, leaving the syringe visually indistinguishable from one that had been filled with nitrone 1b only. No optical signature of a propagating reaction-diffusion front is observed in this case. Selected images were processed further (see Supporting Information) to compute the profile (Figure 4b) of the propagating front at a sequence of time points. These data clearly show the progression of the reaction-diffusion front mediated by replicator 3b along the syringe. In many cases, reaction-diffusion fronts propagate at constant linear or radial velocity. However, in this case, the progression of the front slows and will eventually stall as nitrone 1b and maleimide 2 are depleted throughout syringe as a result of the background rate of the cycloaddition reaction forming 3b being significant on the timescale of the experiment.

Here, we have described the first example of a propagating reaction-diffusion front that is initiated and driven by a synthetic replicator. The work reported here represents a proof-of-principle. The successful implementation of a replicator-driven reaction-diffusion front, mediated by an autocatalytic replicator of defined structure and with specific interactions and catalytic relationships with other similar replicators, opens up a number of exciting possibilities. This reaction format will allow us to explore networks of replicators under conditions and outcomes that lie far from the constraints imposed by well-stirred batch reactors. These studies are currently underway in our laboratory.

ASSOCIATED CONTENT

Electronic Supplementary Information (ESI) available: experimental procedures, details of kinetic measurements and fitting, computational modeling of UV spectra, details of UV-Vis and fluorescence analyses and methods and analyses for the reaction-diffusion experiments.

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Replicator-initiated propagating reaction-diffusion front
(a) 

\[
\begin{align*}
\text{1a; 3a: } R &= H \\
\text{1b; 3b: } R &= \begin{array}{c}
\text{phenyl}
\end{array}
\end{align*}
\]
Propagation of an autocatalytic reaction under reaction-diffusion conditions