

# Cooperative Hybridization of Oligonucleotides

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Supporting Information

# ABSTRACT:

Nucleic acids have been demonstrated to be versatile nanoscale engineering materials with the construction of dynamic DNA structures, motors, and circuits. These constructions generally rely on the clever use and integration of relatively few reaction mechanisms and design primitives. Here, cooperative hybridization is introduced as a mechanism in which two oligonucleotides of independent sequence can stoichiometrically, simultaneously, and cooperatively hybridize to a DNA complex. Cooperative hybridization is rigorously characterized and modeled and is shown to implement digital concentration comparison with amplification, as well as digital Boolean logic. These designs, based on cooperative hybridization, excel in being robust to impurities and not requiring oligonucleotide purification.

# **■ INTRODUCTION**

The precise temporal and spatial control of molecules is a fundamental goal of both synthetic biology and nanotechnology and is essential for building reliable nanoscale structures and devices. Nucleic acids, by virtue of their well-understood hybridization thermodynamics and kinetics, exponential information content and 0.4 nm addressability, and economy of synthesis and preparation, have emerged as a leading material for nanoscale engineering. Furthermore, the biological relevance of nucleic acids and the ease of coupling nucleic acids to other materials, such as proteins and carbon nanotubes, facilitate the use of nucleic acids both as synthetic biomaterials and as scaffolds for other nanotechnological applications.

Although the first generation of DNA nanotechnology research has focused on the self-assembly of static DNA nanostructures, <sup>14–18</sup> recent works in the field have also expanded into the realm of constructing dynamic nucleic acid devices, in which nucleic acid nanostructures conditionally and programmably reconfigure in solution. <sup>8,9</sup> Examples include cascaded logical and amplification circuits, <sup>19–27</sup> DNA origami boxes that close and open, <sup>28</sup> molecular walkers that traverse predefined landscapes, <sup>29–32</sup> controlled rotating DNA frameworks, <sup>33–35</sup> and chain reaction DNA motors and dendrimers. <sup>36–38</sup>

While some of these constructions relied on functional nucleic acid molecules with innate catalytic activity (known as ribozymes and deoxyribozymes), <sup>19–22,29,30</sup> many others were constructed using purely rational design approaches, based on the well-characterized thermodynamic <sup>1,39</sup> and kinetic <sup>40,41</sup> properties of DNA

hybridization, branch migration, and dissociation processes. The latter group generally relies the clever and repeated use of a simple but reliable mechanism, known as toehold-mediated strand displacement, in which short, single-stranded domains on different DNA molecules hybridize to colocalize the molecules, enabling subsequent branch migration. <sup>9,33</sup> To further expand the scope of functions achievable with dynamic DNA nanotechnology, it is necessary to develop other molecular mechanisms that afford functionality that toehold-mediated strand displacement cannot achieve.

Here, a generalized method for implementing cooperative hybridization is presented, in which two oligonucleotides (targets) of independent sequence simultaneously and cooperatively bind to a designed two-stranded complex. The complex allows the two targets to stoichiometrically react and colocalize with each other, much in the same way that two complementary oligonucleotides hybridize to each other.

In contrast to toehold-mediated strand displacement reaction networks, which primarily rely on sequential hybridization and branch migration events, cooperative hybridization enables parallel hybridization and branch migration events to occur. In addition to enabling networks such as amplified digital detection of over- and underexpression relative to a threshold, cooperative hybridization also offers the practical advantage of being robust to synthesis impurities and a background of unrelated nucleic acids and thus is likely to be a useful tool for engineering dynamic DNA nanotechnological devices.

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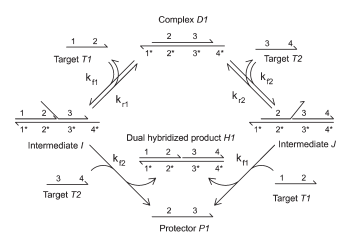


Figure 1. Cooperative hybridization mechanism. Two nucleic acid molecules of interest, T1 and T2, cooperatively hybridize to two-stranded complex D1 (composed of upper strand P1 and lower strand L1). T1 and T2 individually bind to D1 in a reversible manner, but their simultaneous binding releases protector strand P1 and dual-hybridized product H1, rendering the reaction irreversible. DNA strands are represented as directional lines, with the hook denoting the 3' end. Each strand is subdivided into domains, continuous nucleotides that act as a unit in hybridization, branch migration, or dissociation. Domains are represented by numbers, and starred domains denote complements of the unstarred domains (e.g., 2\* is complementary to 2). The  $T1+D1 \rightarrow I$  and  $T1+J \rightarrow P1+H1$  reactions are assumed to have identical rate constants because these reactions are initiated by identical single-stranded toeholds, and similarly for the  $T2+D1 \rightarrow J$  and T2+I→P1+H1 reactions. Previous research has shown that the rate constant of strand displacement reactions are primarily determined by the standard free energy of toehold hybridization.

# **■ COOPERATIVE HYBRIDIZATION MECHANISM**

The cooperative hybridization mechanism is shown in Figure 1. Two target strands, T1 and T2, hybridize simultaneously to two-stranded complex D1 only when both are present. There are two parallel pathways for this reaction. In one, T1 first binds to D1 form intermediate I, which then reacts with T2 to form H1 and release product strand P1. In the other pathway, T2 binds first to D1 to form intermediate I. Individually, the hybridization of T1 or T2 to D1 is reversible and thermodynamically unfavorable; the release of product P1 upon the simultaneous hybridization of T1 and T2 facilitates the net reaction.

The cooperative hybridization mechanism can be expressed as the following reactions:

$$T1 + D1 \rightleftharpoons I$$

$$T2 + D1 \rightleftharpoons J$$

$$T2 + I \rightarrow P1 + H1$$

$$T1 + J \rightarrow P1 + H1$$

The net reaction of the system is thus

$$T1 + T2 + D1 \rightarrow P1 + H1$$

There are three prominent features of this reaction. First, the net reaction possesses different numbers of reactants and products, making the equilibrium distribution of the reactants and products concentration dependent. At low concentrations, entropy is a larger factor, and reactants will exist at higher concentration at equilib-

rium, while the opposite is true at high concentrations. At operational conditions, the products are designed to be predominant.

Second, at operational concentrations where the products are favored, the equilibrium concentration of P1 will be roughly the minimum of the initial concentrations of T1, T2, and D1. As P1 is a strand with a different sequence than both T1 and T2, it can potentially participate in downstream reactions that T1 and T2 cannot. Thus, the cooperative hybridization mechanism not only allows the two target oligonucleotides to be colocalized but also releases a third oligonucleotide to signal the completion of the cooperative hybridization reaction.

Third, because the individual reaction of T1 or T2 with D1 is thermodynamically unfavorable, at operational concentrations very little of T1 or T2 is sequestered in I or J if only one of T1 or T2 is present. The equilibrium concentration of free T1 or T2 in this case would be near the total concentration of T1 or T2. These properties of the cooperative hybridization process allow a variety of useful dynamic nucleic acid devices and circuits.

# ■ MATERIALS AND METHODS

**DNA Sequences and Design.** The conceptual design of the system, such as the domain lengths and relative binding strengths, was based on the expected operational concentration and the desired binding fraction of individual targets in the absence of the second target. In particular, domains 1 and 4 were designed so that, individually, their hybridization energies were insufficient to overcome the entropic loss of colocalizing one additional molecule of DNA, but collectively, they do drive the reaction forward. These considerations are discussed in mathematical detail in Supporting Information, S1.

The sequences were designed subsequently using domain-based sequence design software. The domains (shown in Table 1) possess minimal secondary structure and crosstalk (binding between unrelated domains): NUPACK (http://www.nupack.org/) calculates there to be no more than four paired bases between any pair of strands at 25 °C, even at 1  $\mu$ M concentration. Furthermore, the minimum free energy states of every individual strand was completely unstructured ( $\Delta G^{\circ}=0$ ). Thus, the domains we use can be approximated as structure-free. Substantial secondary structure is known to slow down branch migration and interfere with hybridization.

**Standard Free Energy Calculation.** The standard free energies of complexes are needed in order to calculate the standard free energy of reactions, which in turn can be used to generate equilibrium and rate constants. NUPACK<sup>39</sup> was used to calculate the standard free energies of DNA complexes. NUPACK uses a number of different parameters in its calculations; the values used are detailed and justified below.

Temperature was set to 25 °C, as that was the temperature at which experiments were performed. Salt concentration was set to 0.05 M Na $^+$  and 0.0115 M Mg $^{2+}$ . In actuality, the experimental concentration of Na $^+$  is 0.002 M, but 0.05 M Na $^+$  was the lowest NUPACK allowed. However, since Mg $^{2+}$  acts as the main counterion, it is likely that this difference does not significantly change the standard free energies.

The "dangles" parameter was set to ALL, so that dangles energies are incorporated for all bases flanking duplexes, regardless of whether it is paired. This is necessary because, by default, NUPACK does not incorporate the thermodynamics of coaxial stacks (such as at the nick in *I, J,* and *H*1). The reason NUPACK does not by default include coaxial stacking thermodynamics is that they are still not well-understood: Pyshnyi et al. 44 report that the energetics of coaxial stacking near a nick depends significantly on the nearest neighbors (bases one away from the nick). Setting dangles = ALL allows partial compensation of the energetics at the nicks.

The standard free energies of complexes determine the standard free energies of reactions, which in turn can be used to calculate the equilibrium constant of the reaction via  $\Delta G^{\circ} = -RT \ln(K_{\rm eq})$ . For a reaction with different numbers of reactants and products,  $K_{\rm eq}$  is not unitless, and

Table 1. Domain and Strand Sequences<sup>a</sup>

domain	sequence	length (nt)
1	5'-CATCACTA-3'	8
2 = 2a:2b:2c	5'-CTATCATCACACATCTAT-3'	18
2a	5'-CTATCAT-3'	7
2b	5'-CACACAT-3'	7
2c	5'-CTAT-3'	4
3 = 3a:3b	5'-ACAACCACTTACTTCTTC-3'	18
3a	5'-ACAACCACTTACTT-3'	14
3b	3'-CTTC-3'	4
4	5'-ATCTATCC-3'	8
5	5'-CTATCAT-3'	7
6	5'-CACACAT-3'	7
7	5'-CTATACAACCACTTACTT-3'	18
8	5'-CTTC-3'	4
9	5'-GCCATCAGAACTTAACCT-3'	18
10	5'-AACTC-3'	5
11	5'-CTTTCCTACA-3'	10
12a	5'-CCTACGTCTC-3'	10
12b	5'-CAACTAA-3'	7
12c	5'-CTTACGG-3'	7
13	5'-CCCTC-3'	5
strand	domain composition	length (nt
T1	1 2	26
T2	3 4	26
P1	2 3	36
L1	4* 3* 2* 1*	52
F	2c 3a	18
RL	3a* 2c* 2b*	25
Т3	5 6 7 8	36
T4	9 10 ROX	23
P2	7 9	36

<sup>&</sup>lt;sup>a</sup> The sequences of the starred complement domains are determined by the corresponding unstarred domains. For example, the 10 domain has sequence 5'-AACTC-3', so 10\* has sequence 5'-GAGTT-3'. Strands are shown as concatenations of domains, listed from 5' to 3' end.

RQ 10\* 9\* 7\* 6\*

L2

its value will depend on its unit. Thus, the  $\Delta G^{\circ}$  of a reaction may depend on the units used for expressing  $K_{\rm eq}$ . For example,  $K_{\rm eq}=1$  M is identical to saying  $K_{\rm eq}=1000$  mM, but the former yields  $\Delta G^{\circ}=0$  kcal/mol, while the latter yields  $\Delta G^{\circ}=-4.1$  kcal/mol at 25 °C.

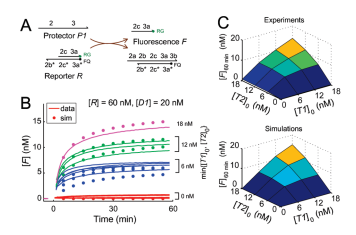
NUPACK reports the  $\Delta G^{\circ}$  values of complexes based on the equilibrium constant of the complex's formation, calculated in mole fraction. Because the concentration of all nucleic acid molecules is negligible compared to that of water ((1000 g/L)/(18 g/mol) = 55 M), the molarity of the oligonucleotides is roughly 1/55th that of their mole fractions. To convert the NUPACK reported  $\Delta G^{\circ}$  into one that yields  $K_{\rm eq}$  expressed in molar, we add a corrective  $(n-1)RT\ln(55)$  term, where n denotes the number of strands the DNA molecule possesses (i.e., +2.38 kcal/mol for every strand in excess of 1).

Intermediates I and J represent the three-stranded complexes on which branch migration is possible (domain 2 for I and domain 3 for J). Consequently, I and J each correspond to 19 isoenergetic branch migration states ( $I_0$  through  $I_{18}$  and  $J_0$  through  $J_{18}$ , respectively). Each

Table 2. Composition and Standard Free Energies of Complexes Shown in Figures 1 and  $2^a$ 

complex	strand composition	calcd $\Delta G^\circ$ (kcal/mol)
D1	P1, L1	-49.27 + 2.38 = -46.9
I	P1, L1, T1	-62.15 + 2(2.38) - 1.75 = -59.1
J	P1, L1, T2	-61.51 + 2(2.38) - 1.75 = -58.5
H1	L1, T1, T2	-70.69 + 2(2.38) = -65.9

 $^a$  The standard free energies of these complexes were calculated using NUPACK,  $^{39}$  using the parameters 25 °C, 0.05 M Na $^+$ , 0.0115 M Mg $^{2+}$ , dangles = ALL. The standard free energies of all individual strands were 0 kcal/mol (completely unstructured). Because NUPACK yields  $\Delta G^\circ$  values that were calculated for mole fraction rather than molar, a corrective  $RT\ln([\mathrm{H_2O}])=+2.38$  kcal/mol term must be added for every strand in excess of 1. The intermediate states I and J each correspond to 19 isoenergetic branch migration states; the  $-RT\ln(19)=-1.75$  kcal/mol term corrects for this state multiplicity.



**Figure 2.** Characterizing and modeling cooperative hybridization. (A) The kinetics of the cooperative hybridization reaction are assayed through the use of a fluorescence reporter complex R (comprised of F and RL). P1 reacts stoichiometrically with R to release fluorophore-labeled strand F. RG denotes Rhodamine Green, and FQ denotes the Iowa Black Fluorescence Quencher. (B) Kinetics of cooperative hybridization. R and D1 were present in solution initially, and various quantities of T1 and T2 were introduced at time  $t \approx 0$ . Because D1 is in excess of both T1 and T2, both the kinetics and the final equilibrium fluorescence value depend on the minimum of the concentrations of T1 and T2. The dotted lines show simulation results using experimentally measured rate constants (see Table 3, Supporting Information, S3 and Figure S1). (C) Summary of experimental and simulations results, with fluorescence plotted against the concentrations of T1 and T2. This figure shows that the release of P1 (and subsequently F) is contingent upon the simultaneous presence of T1 and T2.

of these states has a  $\Delta G^{\circ}$  calculated by NUPACK to satisfy the equilibrium constant in one of the following reactions:

$$T1 + D1 \rightleftharpoons I_0 \rightleftharpoons I_1 \cdots I_{17} \rightleftharpoons I_{18}$$

$$T2 + D1 \rightleftharpoons J_0 \rightleftharpoons J_1 \cdots J_{17} \rightleftharpoons J_{18}$$

The standard free energies of the amalgamate states I and J are defined so as to preserve the equilibrium concentrations of D1 and T1 or T2. To do so,  $-RT\ln(19) = -1.75$  kcal/mol is added to the  $\Delta G^{\circ}$  of a single  $I_0$  or  $J_0$  state, respectively, to derive the  $\Delta G^{\circ}$  of I or J.

Table 2 shows the calculated free energies of the complexes.

**Annealing.** All annealing processes were performed with an Eppendorf Mastercycler Gradient thermocycler. The samples were brought down from 95 to 20  $^{\circ}$ C at a constant rate over the course of 75 min.

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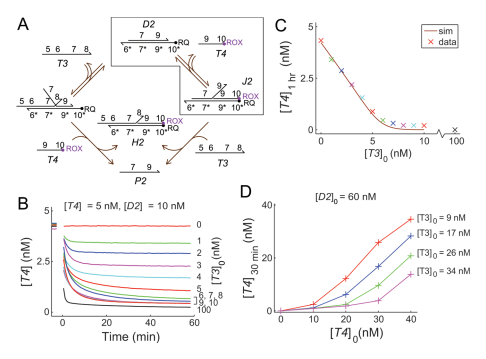


Figure 3. Demonstration of cooperativity. (A) T3 and T4 cooperatively hybridize to D2. T4 is fluorophore-labeled, and the bottom strand of D2 is quencher-labeled. When T4 is hybridized to D2, the fluorophore is colocalized to the quencher and fluorescence decreases. In the absence of T3, T4 should not significantly bind to D2 at the operational conditions, so fluorescence should be high. The box shows the species present before T3 is added. (B) Experimental results support the proposed mechanism. D2 and T4 are initially present, and fluorescence is similar to that of T4 alone. As various amounts of T3 are introduced into the reaction at  $t \approx 0$ , the fluorescence decreases. (C) Summary of the results in (B). Fluorescence after 1 h follows a kinked line, with the kink roughly at [T3] = 5 nM; errors in the concentration of the stock solutions of T3 and T4 likely contribute to the kink deviating from 5 nM. In fact, the cooperative hybridization mechanism can be used as a method of quantitating nucleic acid concentrations given a known standard of independent sequence (see also Supporting Information, S4 and Figure S3). Simulation traces shows the predicted results according to our model, fitting the T3 concentration to be 14% lower than as measured by absorbance. (D) The equilibrium concentration of free T4 (which is roughly linear to fluorescence) is the difference between the initial concentrations of T4 and T3. Data displayed shows the fluorescence after 30 min, as a function of T3 and T4 concentration. Fluorescence appears to be roughly linear in the concentration excess of T4 over T3. This figure shows that the individual binding of T3 or T4 to D2 is reversible and not favorable at equilibrium, and that the hybridization mechanism is cooperative as designed.

Complex Purification. DNA oligonucleotides used in this study were purchased from Integrated DNA Technologies (IDT). Where applicable, fluorophores were attached by IDT as well. Concentrations were determined from the measured absorbance at 260 nm using an Eppendorf Biophotometer and calculated extinction coefficients. 45,46

Complex D1 used in Figure 2 was purified by non-denaturing (ND) polyacrylamide gel electrophoresis (PAGE) to ensure proper stoichiometry as follows: The P1 and L1 strands were prepared with nominally correct stoichiometry at 20  $\mu$ M and annealed. The samples were then run on 12% ND PAGE at 180 V for 6 h.

The acrylamide (19:1 acrylamide:bis) was diluted from 40% acrylamide stock (Ambion). ND loading dye containing xylene cyanol FF in 50% glycerol was added to all samples, achieving a final gycerol concentration of 10% by volume. Gels were run at 25  $^{\circ}$ C using a Novex chamber with an external temperature bath.

The proper D1 band was cut out and eluted in 2 mL of TE/Mg<sup>2+</sup> buffer for 2 days. Purified complexes were quantitated by 260 nm absorbance measurement and calculated extinction coefficients. Yield was approximately 50%.

Notably, only the complex D1 used for experiments in Figure 2 was purified; the D2 used in Figure 3 and the D1 used in Figures 4—6 were not purified.

**Buffer Conditions.** DNA oligonucleotides were stored in TE buffer (10 mM Tris·HCl pH-balanced to 8.0, with 1 mM EDTA·Na<sub>2</sub>, purchased as 100x stock from Sigma-Aldrich) at 4 °C. Directly preceding experiments, TE buffer with 62.5 mM MgCl<sub>2</sub> was added at a 1:4 ratio to the sample, achieving a final MgCl<sub>2</sub> concentration of 12.5 mM (of which 1 mM is bound

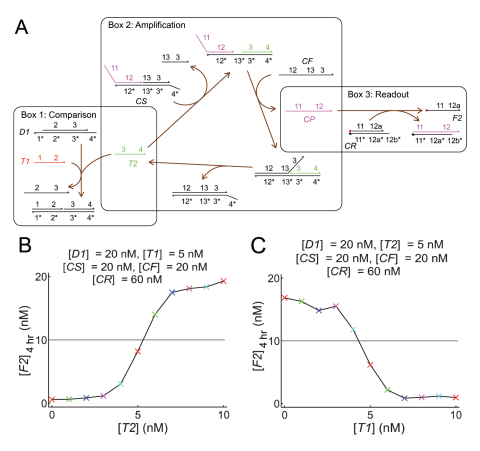
to EDTA). This buffer is henceforth known as "TE/Mg $^{2+}$ " buffer. All experiments and purifications were performed at 25  $\pm$  0.5 °C, with temperature controlled using an external temperature bath.

**Spectrofluorimetry Studies.** Spectrofluorimetry studies were done using a SPEX Fluorolog-3 (Horiba) with 1.6 mL 119-004F synthetic quartz cells (Hellma). For the experiments shown in Figures 3, 5D, and 6, excitations were at 588 nm, while emissions were at 602 nm (optimal signal for ROX fluorophore). For the experiments shown in Figures 2, 4, and 5B, excitations were at 510 nm, while emissions were at 531 nm (optimal signal for Rhodamine Green fluorophore). In all spectrofluorimetry experiments, the total reaction volume was 1.5 mL. For Figures 3B,C, 5E, 6, and S2A—C, 4 nm band-pass slits were used for both excitation and emission monochromators. For all other experiments, 2 nm slits were used. In all experiments, data points were collected with an integration time of 10 s for every 60 s time point.

Prior to each experiment, all cuvettes were cleaned thoroughly: each cuvette was washed 15 times in distilled water, once in 70% ethanol, another five times in distilled water, and finally once more in 70% ethanol.

For the slit size, concentrations, and times chosen, no measurable photobleaching was observed. All experimental results were within the linear regime of the spectrofluorimeter detector, according to the specification sheets provided by the manufacturer.

**Fluorescence Normalization.** Fluorescence is normalized so that 1 normalized unit (n.u.) of fluorescence corresponds to 1 nM of an unquenched fluorophore-labeled strand (F in Figures 2 and 5B, T4 in Figures 3, 5D, and 6, and F2 in Figure 4). This normalization is based on the fluorescence levels of annealed samples: A negative control with  $\lceil R \rceil$ 



**Figure 4.** Application: amplified digital detection of over- and under-expression relative to a threshold. (A) Schematic. The detection circuit is divided into three modular components, shown as boxes. For over-expression detection, T1 serves as the threshold and is present at known concentration. If the amount of T2 exceeds that of T1, then the excess T2 catalyzes the release of CP from CS, which in turn reacts stoichiometrically with fluorescent reporter CR to release fluorophore-labeled strand F2 (see also ref 25). For under-expression detection, T2 serves as the threshold with known concentration and will trigger the catalytic pathway unless excess T1 is added. (B) Over-expression detection. Fluorescence increases as  $[T2]_0$  exceeds the threshold set by  $[T1]_0$ . T2 is pre-reacted with D1 and T1 for T1 min, and then T1 for T1 min, and then T1 for T1 min, and then T1 for T1 for T1 min, and the catalytic substrate and reporter species. (C) Under-expression detection. Fluorescence decreases as  $[T1]_0$  exceeds the threshold set by  $[T2]_0$ .

= 30 nM yielded the fluorescence of the quenched F strand (Rhodamine Green), and a positive control with [R] = 30 nM and [P1] = 10 nM yielded the fluorescence of the unquenched F strand. Similarly, a negative control with [T4] = 5 nM and [L2] = 10 nM yielded the fluorescence of the quenched T4 strand (ROX), and a positive control with [T4] = 5 nM yielded the fluorescence of the unquenched T4 strand.

Day-to-day and sample-to-sample variations in fluorescence are estimated to be less than 5%.

Ordinary Differential Equation (ODE) Simulations. Reaction simulations were run in Matlab using the "stiff" ode23s solver because of the difference in time scales of bimolecular and unimolecular reactions. The relative tolerance of the solver was set to  $10^{-4}$ , and the absolute tolerance of the solver was set to  $10^{-30}$  M. Supporting Information, S2 shows sample code for simulating the system shown in Figure 2.

**Parameter Fitting.** Rate constants were fitted to experimental data using the "fminunc" function in Matlab to minimize the error between experimental data and the reaction model. The error between the data and the simulation is calculated as

Error = 
$$\sum_{t, \text{traces}} (F_{d}(t) - F_{m}(t))^{2}$$

where  $F_d(t)$  is the fluorescence value of the data at time t, and  $F_m(t)$  is the fluorescence value predicted by the ODE model at time t. Sample code for fitting the rate constants is shown in Supporting Information, S2.

The confidence intervals on the values of the rate constants were generated as the values of the rate constant at which the error score is double that of the minimum. All other rate constants were kept constant at their best-fit values when generating the confidence interval on each rate constant.

# **■ RESULTS**

Characterizing Cooperative Hybridization. To demonstrate that the cooperative hybridization mechanism functions as intended, two different properties must be independently verified: first that T1 and T2 when simultaneously present can cooperatively hybridize to D1, and second that T1 in the absence of T2 or T2 in the absence of T1 is insufficient to stably bind D1.

Figure 2 shows the kinetic behavior of the cooperative hybridization system through the use of a fluorescent reporter complex R that reacts stoichiometrically with P1. The release of P1 and the subsequent fluorescence increase is seen to vary linearly with the minimum of the initial concentrations of T1 and T2, supporting the proposed mechanism.

Also shown in Figure 2B,C are simulation results, using individually measured rate constants (see Supporting Information, S3 and Figure S1). The reactions simulated and their relevant rate constants are summarized in Table 3. The agreement between

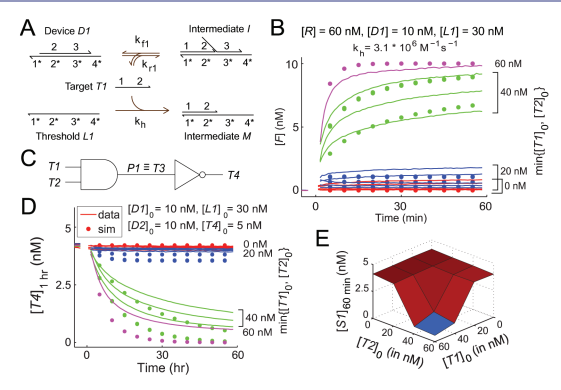
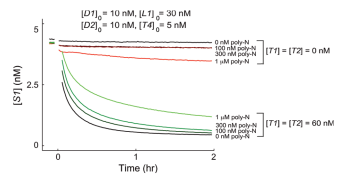


Figure 5. Application: cascaded nucleic acid logic. (A) Thresholding using the bottom strand L1 of D1. Targets T1 and T2 preferentially bind L1 over D1 because L1 offers a longer single-stranded toehold for initiating hybridization and because individual hybridization is irreversible. (B) Experimental and simulation results of cooperative hybridization with a threshold. The fluorescence increases quickly when the concentrations of both targets T1 and T2 exceed that of the threshold L1. (C) The P1 species from Figures 1 and 2 was designed to be identical in sequence to T3 in Figure 3. Consequently, the logical AND and the logical NOT gates can be directly cascaded to form a network exhibiting NAND logic. (D) Experimental data and simulations on the logical NAND cascaded network. D1, L1, D2, and T4 were present in solution initially, T1 and T2 were added at time  $t \approx 0$ . No new parameters were fit to the data shown in this figure; rate constants simulated were fitted in previous figures. (E) 3D summary of the experimental results in part (D).



**Figure 6.** Robustness to background molecules. Various amounts of poly-N mix (50 nt oligonucleotides with a random distribution of G, C, A, and T bases at every position) were added the NAND reaction network. D1, L1, D2, and T4 were present in solution initially; T1, T2, and poly-N were added at time  $t \approx 0$ .

ODE simulations and experimental data further supports that the cooperative hybridization mechanism functions as designed: the consumption of T1, the consumption of T2, and the production of P1 are all simultaneous and stoichiometric with respect to each other. The small quantitative differences between simulations and experiments are likely due to oligonucleotide impurities (see Supporting Information, S6 and Figure S5) and undocumented side reactions (such as  $T2 + R \rightarrow F$ ).

Figure 3 demonstrates that the two DNA oligonucleotide targets do not unilaterally bind irreversibly to the two-stranded complex. Here, different sequences are used, with T3 and T4

cooperatively hybridizing to complex *D*2. *T*4 is labeled with the ROX fluorophore, and the bottom strand *L*2 of the *D*2 complex is labeled with the Iowa Black Red Quencher. Thus, when *T*4 is hybridized to *D*2, either temporarily or permanently, the observed fluorescence is decreased. This allows a direct assay of the instantaneous concentration of free *T*4.

The other oligonucleotide target *T*3 possesses both 5' and 3' overhangs that serve no function for the cooperative hybridization mechanism. The existence of these overhangs demonstrates that the cooperative hybridization mechanism can be used to target a subsequence of a longer nucleic acid, such as an mRNA.

Figure 3B shows the results of this system. The fluorescence of solution in the absence of T3 is between 4 and 4.5 nM, while  $[T4]_0 = 5$  nM. A small fraction of the T4 is associated to D2 at equilibrium. When an amount of T3 is added to the system, the fluorescence (concentration of free T4) re-equilibrates, consistent with our understanding of the cooperative hybridization mechanism.

The stoichiometric consumption of the two single-stranded reactants in cooperative hybridization can be interpreted as analog subtraction. If initially  $[T4]_0 > [T3]_0$ , then the equilibrium concentration of free T4 will be  $[T4]_\infty \approx [T4]_0 - [T3]_0$ . This property can be used to compare the concentrations of oligonucleotides of known sequences. For example, T4 can act as a well-characterized standard, and a different D2 complex can be constructed for each nucleic acid molecule T3 of interest. In a solution of a known quantity of T4 and an excess of D2, the concentration of T3 can be inferred from the equilibrium fluorescence (see Supporting Information, S4 and Figure S3). Because the same standard T4 can be used to quantitate a variety of

Table 3. Reaction Kinetics and Thermodynamics

Reactions and Rate Constant	s Simulated in Figure 2B,C"
$T1+D1 \stackrel{k_{f1}}{\underset{\stackrel{\leftarrow}{\longrightarrow}}{\rightleftharpoons}} I$	$k_{\rm f1} = (2.1 \pm 0.7) \times 10^6 \mathrm{M}^{-1}\mathrm{s}^{-1}$
$T2 + D1 \stackrel{\tau_1}{\rightleftharpoons} J$	$k_{\rm f2} = (1.4 \pm 0.2) \times 10^6 \mathrm{M}^{-1}\mathrm{s}^{-1}$

$$T2 + I \stackrel{k_{12}}{\rightleftharpoons} P1 + H1$$
  $k_{r1} = 0.54 \pm 0.42 \text{ s}^{-1}$ 

$$T1 + J \xrightarrow{k_{\text{rp}}} P1 + H1$$
  $k_{\text{r2}} = 0.39 \pm 0.27 \text{ s}^{-1}$   $P1 + R \xrightarrow{k_{\text{rp}}} F + Fw$   $k_{\text{rep}} = (1.3 \pm 0.5) \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ 

Standard Free Energies of the Reactions Simulated in Figure 2B,C<sup>b</sup> calcd  $\Delta G^{\circ}$  (kcal/mol) fitted  $\Delta G^{\circ}$  (kcal/mol)

$T1 + D1 \underset{k_1}{\overset{k_{f1}}{\rightleftharpoons}} I$	-12.2	-9.00
$T2 + D1 \stackrel{ii}{\underset{k_{12}}{\rightleftharpoons}} J$	-11.6	-8.95
$T2+I \stackrel{k_{f2}}{\longrightarrow} P1+H1$	-6.8	N/A
$T1+J \stackrel{k_{\rm f1}}{\longrightarrow} P1+H1$	<b>-7.4</b>	N/A

<sup>a</sup> The simulations use the best-fit value of the rate constants; the errors bars on the rate constants are calculated as described in Materials and Methods, Parameter Fitting.  $^b$  Calculated  $\Delta G^\circ$  (25  $^\circ$ C) denotes the value obtained from using the NUPACK predicted standard free energies of the complexes, and the fitted  $\Delta G^{\circ}$  denotes the value obtained from  $\Delta G^{\circ} = -RT \ln(k_{\rm f}/k_{\rm r})$  The fitted  $\Delta G^{\circ}$  of the latter two reactions cannot be calculated because the reverse reaction was too slow to observe: according to the calculated  $\Delta G^{\circ}$ , the reverse rate constant would be around 10 M<sup>-1</sup> s<sup>-1</sup>, corresponding to a time scale of about 3 months at an operational concentration of 10 nM.

Table 4. Reactions Simulated in Figure 3B,C<sup>a</sup>

reactions	rate constants
$T3 + D2 \stackrel{k_{f3}}{=} I2$	$k_{\rm f3} = 4 \times 10^6 \mathrm{M}^{-1} \mathrm{s}^{-1}$
$T4 + D2 \stackrel{\sim}{\underset{k_4}{\rightleftharpoons}} J2$	$k_{\rm f4} = 4 \times 10^6 \mathrm{M}^{-1} \mathrm{s}^{-1}$
$T4+I2 \stackrel{iq}{\longrightarrow} P2+H2$	$k_{\rm r3} = 0.3 \; \rm s^{-1}$
$T3+J2 \stackrel{k_{\rm f4}}{\longrightarrow} P2+H2$	$k_{\rm r4} = 0.2 \; \rm s^{-1}$

<sup>a</sup>The data displayed in Figure 3 underconstrain the rate constants. Shown here are one set of rate constant values that generated reasonably good agreement between ODE simulations and experimental data. Similar qualities of fit can be attained by co-varying the four rate constants.

different oligonucleotides, quantitation using cooperative hybridization may yield advantages over methods based on molecular beacons<sup>47</sup> because of the reduced need for calibration.

From the data shown in Figure 3B, the concentrations of T3 and T4 are implied to differ from their nominal values calculated from absorbance at 260 nm. Arbitrarily assuming that the concentration of T4 is accurate, the true concentration of T3 is inferred to be 14% lower than given. This 14% difference may arise due to errors in extinction coefficients, sample impurities that yield absorbance at 260 nm, and/or truncated oligonucleotides that cannot undergo cooperative hybridization. Figure 3C shows simulation results assuming that the concentrations of T3 are 14% lower than listed, and Figure 3D shows the equilibrium concentrations of T4 based on a variety of initial concentrations of *T*3 and *T*4 (with the corrected concentration of *T*3).

Table 4 shows the reactions simulated to generate the simulation traces shown in Figure 3C. Five different parameters were fitted to the data in Figure 3C (the four rate constants and the concentration of T3). Given the limited data on this system, the fit was underconstrained, and it is likely that many different sets of rate constant values would have yielded fits of similar quality.

The length of domain 6 is 7 nt, rather than 8 nt as in domains 1 and 4, because of the thermodynamically stabilizing effect of the 5' dangle. The length of domain 10 is 5 nt, rather than 7 or 8, because of the stabilizing effects of fluorophore-quencher binding on the thermodynamics of DNA hybridization.<sup>48</sup> Our own experiments suggest that the interaction between ROX and Iowa Black Red Quencher is similar to that of 4-5 base pairs binding (data not shown).

Reaction Networks Involving Cooperative Hybridization. Cooperative hybridization is a modular dynamic DNA nanotechnology component that can be integrated with other systems to form reaction networks with desired properties. In Figure 4, cooperative hybridization is cascaded with a DNA hybridizationbased catalyst system<sup>25</sup> and a fluorescent reporter to yield a network that performs digital concentration comparison of two different oligonucleotides. This could potentially be used for detecting over-expression and under-expression of nucleic acids relative to certain thresholds.

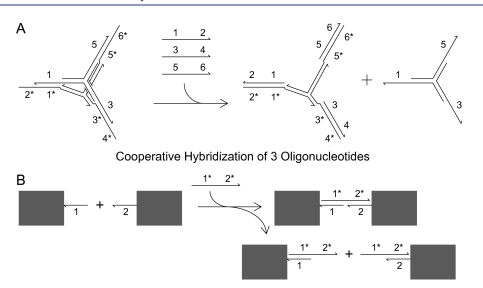
In this reaction network, T2 serves as the catalyst for releasing CP from multistranded complex CS. Because T2 cooperatively hybridizes to D1 with T1, if the concentration of T1 exceeds that of T2, then little to no free T2 will exist, and CP will not be released. In contrast, if [T2] exceeds [T1], even if only by a little, then it catalytically releases CP, and the final fluorescence value will be determined by the concentration of precursor CS initially present in solution. Thus, the thresholded linear response shown in Figure 3C is converted to the sigmoidal thresholded response shown in Figure 4B,C. Alternative methods of digitally comparing nucleic acid concentrations using cooperative hybridization also exist (see Supporting Information, S5 and Figure S4).

In addition to providing a sigmoidal thresholded response, this reaction network also amplifies the fluorescence signal relative to the concentrations of the targets. In Figure 4B,C, the concentrations of T1 and T2 were both less than 10 nM, yet the final fluorescence corresponded to up to 20 nM of released fluorophore-labeled strands, demonstrating thresholded amplification using purely DNA components. Further amplification can be achieved through multilayer catalyst cascades or feedback systems. 25,49

Logical Processing Using Cooperative Hybridization. The properties of cooperative hybridization can be interpreted as implementations of Boolean logic. For example, in Figure 1, P1 can be considered the output of a logical AND gate, with T1 and T2 being the inputs. Alternatively, in Figure 3, T4 can be considered the output of a logical NOT gate, with T3 being the input.

One key feature of Boolean logic is digital abstraction, wherein analog values are converted to digital ones through sigmoidal thresholding and amplification, both of which have been demonstrated in Figure 4. However, a simpler method of thresholding can be achieved using excess L1, the bottom strand of D1 (Figure 5A). Targets T1 and T2 preferentially bind L1 over D1 because L1 offers a longer single-stranded toehold for initiating hybridization and because individual hybridization is irreversible. Figure 5B shows that when either [T1] or [T2] is less than [L1], little product P1 is released; this is consistent with the assumption that *T*1 and *T*2 preferentially bind to *L*1.

Different cooperative hybridization elements can be cascaded with one another to exhibit more advanced logical functions. Here, the AND gate of Figure 1 is cascaded to the NOT gate of Figure 3 to form a reaction network exhibiting NAND logic (Figure 5C). Ordinarily, this would require one or more strand



Cooperative Hybridization as a Method of Joining Nanostructures

Figure 7. Potential extensions of cooperative hybridization. (A) Proposed method for extending cooperative hybridization to more than two targets. Note that the domain numbers present in this figure do not correspond to the domains shown in previous figures. Here, hybridization of even two targets to the device is reversible; only when all three targets bind does the top three-arm junction get released. (B) Proposed application of cooperative hybridization as a means of reliably joining DNA nanostructures. Joining DNA complexes or nanostructures using a complementary linker can cause unintentionally capping and prevent proper joining (top). In the cooperative hybridization mechanism (bottom), each overhang cannot individually hybridize stably to the linker, and joining can reliably occur.

displacement reactions to stoichiometrically convert P1 into T3.<sup>23</sup> However, in this case T3 was designed to be identical in sequence to P1, so that this stage can be eliminated. Figure 5D,E shows that the NAND network functions as designed: the observed fluorescence decreases only when both [T1] and [T2] exceed the threshold [L1].

**Robustness.** Thus far, all reactions have been tested in isolated solutions, in which no nucleic acids other than those needed were present. In order for the designed systems to function in either a biological setting or a complex synthetic chemical network, however, all designed reactions must be robust to a background of nucleic acids that interact nonspecifically.

Here, the robustness of cascaded cooperative hybridization reaction networks is tested by running the NAND reaction network in a solution of poly-N strands (Figure 6). The poly-N strands are each 50 nt long, with a random (G, C, A, T) base at every position with roughly equal probability. As  $4^{50} > 10^{23}$ , it is likely that every strand in the poly-N mix is different in sequence.

The reaction network functions qualitatively similarly, even in the presence of a  $16\times$  excess of poly-N over T1 and T2, attesting to the robustness and specificity of the cooperative hybridization mechanism. A standard hybridization assay is likely to be more disrupted by the presence of these poly-N strands, due to nonspecific binding interactions. Thus, the cooperative hybridization mechanism provides an additional degree of robustness and specificity over typical nucleic acid hybridization.

**Extensions of Cooperative Hybridization.** Cooperative hybridization is a general design principle for engineering nucleic

acid devices and is by no means limited to those examples detailed in this manuscript. In this section, two extensions of the cooperative hybridization mechanism are proposed.

Cooperativity is a key feature of ensuring specific binding in both biological enzyme function and synthetic biotechnological devices. To construct devices that sense the simultaneous presence/colocalization of many oligonucleotides, increased cooperativity with Hill coefficient higher than 2 is desirable. One potential method of extending the cooperative hybridization mechanism introduced here is shown in Figure 7A. All three target oligonucleotides must be present to drive the hybridization reactions forward; any two of the three targets in isolation is insufficient and leads to only transient binding.

Similar methodology can be used to construct systems requiring four or more cooperative hybridization events. Alternatively, asymmetric branched DNA can also be used to implement three-and four-target cooperative hybridization. In cooperative hybridization of three or more oligonucleotides of independent sequence, geometry needs to be considered in addition to hybridization thermodynamics.

The controlled joining of DNA complexes or nanostructures could allow hierarchical, isothermal assembly of larger DNA nanostructures while preserving nanometer addressability. The use of linker oligonucleotides in a sandwich assaylike process, in which each subunit hybridizes irreversibly to the linker, can cause the subunits to be capped by the linker and preclude proper joining (Figure 7B). In the cooperative hybridization mechanism, each subunit individually cannot stably hybridize to the linker, so this problem is averted.

# DISCUSSION

A cooperative nucleic acid hybridization mechanism was presented and verified in this work. In this mechanism, a two-stranded complex was designed so that two oligonucleotides of independent sequence can cooperatively and simultaneously hybridize to it. The cooperative hybridization mechanism is shown to be robust and modular, smoothly integrating with other dynamic DNA components to form cascaded reaction networks that perform a variety of functions.

The reactions shown here were rationally designed on the basis of the biophysics of nucleic acid hybridization, branch migration, and dissociation. Previous characterization of similar reactions has demonstrated that they function robustly across a wide range of solution salinities and temperatures, <sup>25</sup> as well as in the presence of total RNA and cell lysate. <sup>23,25</sup> Furthermore, similar strand displacement-based RNA devices have been made to assay and regulate gene expression in cells. <sup>50–52</sup> Thus, it is likely that many of these devices can be made to function *in situ* and *in vivo* for assaying and regulating gene expression.

Cooperative hybridization was demonstrated to implement analog subtraction and minimum functions. Both can serve important roles in the analysis and evaluation of biological nucleic acids, in which over- and under-expression of an RNA relative to a standard expression level can be indicative of disease. Integrating these elements with amplification and readout could potentially allow the construction of nucleic acid devices that perform embedded computation within cells, such as determining cell state from microRNA concentrations.

These analog primitives can also serve useful functions in the construction of synthetic circuits for controlling nanoscale self-assembly. In biology, cellular differentiation and development are often guided by the local relative expressions of different genes. Similarly, reaction—diffusion systems combining nucleic acid concentration gradients with concentration comparison circuits could lead to the formation of complex patterns and structures.

By combining cooperative hybridization with thresholding mechanisms, digital logic gates and circuits were constructed. Compared to previous implementations of nucleic acid logic gates and circuits based on strand displacement, <sup>23,26</sup> the major advantage of logic gates based on cooperative hybridization is the ease of sample preparation: the two-stranded complexes used for cooperative hybridization did not require any purification, either at the strand level or at the complex level. Furthermore, these systems are robust to nonspecific oligonucleotides coexisting in solution. These technical advantages reduces the labor needed to set up these circuits, thereby facilitating the construction of more complex reaction networks.

However, like previous implementations, the cooperative hybridization implementation of AND and NOT gates possesses certain limitations. Most notably, cooperative hybridization is a kinetically irreversible process, and logical values, once set, can never be reset. As a concrete example, in the NOT implementation (Figure 3), the *T*4 output strand binds irreversibly to *D*2 and *T*3, and so the output value of the NOT gate can only be changed from ON to OFF, and never from OFF to ON. This practically means that the NOT gate is not dynamic—it cannot, for example, be cascaded to form feedback circuits that require multiple value changes, such as a three-ring oscillator composed of three serially cascaded NOT gates in a feedback loop. Implementation of a dynamic nucleic acid-based logical NOT gate, in which the output

concentration dynamically reflects changes in input concentrations, remains an open problem.

The cooperative hybridization mechanism enables more than just nucleic acid detection, quantitation, and logic; it is a fundamentally new design primitive that can allow simultaneity detection, precise timing control, nonlinear signal responses, and nanostructure joining. Integration of the cooperative hybridization mechanism with functional nucleic acids such as aptamers and ribozymes, are expanded nucleic acid alphabets, bn DNA-directed chemical synthesis, or other nanomaterials can broaden the set of chemistries that can serve as both input and output of an engineered nucleic acid system.

The complexity of natural biochemical circuits enables wondrous behaviors of life such as development, metabolism, and reproduction. It remains an outstanding goal of synthetic biology to rationally design reaction networks that exhibit similar spatial/temporal control of biochemistry; the design, demonstration, and integration of modular nucleic acid systems is one promising approach. The cooperative hybridization mechanism demonstrated in this paper may play an important role in constructing complex synthetic nucleic acid reaction and control networks.

#### ASSOCIATED CONTENT

Supporting Information. DNA sequence design, Matlab code for simulating reactions and fitting rate constants, characterization of individual rate constants, concentration quantitation from Figure 2, alternate mechanism for digitally comparing nucleic acid concentrations, and characterization of oligonucleotide purity using denaturing PAGE. This material is available free of charge via the Internet at http://pubs.acs.org.

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