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## **OPEN** Cooperative control of a DNA origami force sensor

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Biomolecular systems are dependent on a complex interplay of forces. Modern force spectroscopy techniques provide means of interrogating these forces, but they are not optimized for studies in constrained environments as they require attachment to micron-scale probes such as beads or cantilevers. Nanomechanical devices are a promising alternative, but this requires versatile designs that can be tuned to respond to a wide range of forces. We investigate the properties of a nanoscale force sensitive DNA origami device which is highly customizable in geometry, functionalization, and mechanical properties. The device, referred to as the NanoDyn, has a binary (open or closed) response to an applied force by undergoing a reversible structural transition. The transition force is tuned with minor alterations of 1 to 3 DNA oligonucleotides and spans tens of picoNewtons (pN). The DNA oligonucleotide design parameters also strongly influence the efficiency of resetting the initial state, with higher stability devices ( $\gtrsim 10 \text{ pN}$ ) resetting more reliably during repeated force-loading cycles. Finally, we show the opening force is tunable in real time by adding a single DNA oligonucleotide. These results establish the potential of the NanoDyn as a versatile force sensor and provide fundamental insights into how design parameters modulate mechanical and dynamic properties.

Biomolecular functions are often driven by inter- and intramolecular forces. Thus, elucidating the forces within and between biomolecular systems provides critical insight into the mechanisms of their functions<sup>1-3</sup>. Molecular force spectroscopy has been a powerful approach for probing the interactions that are responsible for these forces and providing mechanistic insight into function<sup>4-7</sup>. However, current force spectroscopy techniques have limitations such as challenges with force measurements in constrained environments. For instance, both magnetic and optical tweezers necessitate the use of large (>1  $\mu$ M) beads, which act as handles for applying forces on nanoscale samples<sup>6-9</sup>. Atomic force microscopy requires the sample be attached to a cantilever tip<sup>4,6,10-13</sup>. These methodologies are limited to systems where space is available for the handles, which makes it challenging to implement these approaches within cells<sup>14,15</sup> and nanofluidic devices<sup>16,17</sup>. Nanomechanical devices are a promising alternative approach to probe molecular forces, but this requires versatile device designs that can easily be tuned to respond to a wide range of forces. Here we present the development of a DNA Origami (DO) nanodevice that has the potential to address these challenges, with a focus on establishing simple changes in design parameters that allow versatile tuning of the force response.

DO nanotechnology has significant promise in developing nanodevices for complex functions including drug delivery<sup>18-20</sup>, molecular sensing<sup>21,22</sup>, and probing single molecule dynamics and interactions<sup>23-28</sup>. More specifically, DO has been established as a useful approach for single molecule force sensing, with demonstration of DO devices applying and responding to both tensile and compressive forces<sup>29-32</sup>. Complex and dynamic 3-dimensional DO nanodevices can perform prescribed functions through controlled actuation, making their use precise and reproducible<sup>30,33-35</sup>. DO devices are biocompatible, functionalizable, and on the nanometer (nm) size scale, which are key characteristics that position them to function within complex nanoscale environments. For example, DNA duplexes<sup>36,37</sup> and DO platforms<sup>29</sup> have been successfully implemented to investigate cellular forces by connecting these constructs between a cell and a surface. In the case of DNA duplexes, distinct constructs allow the measurement of different forces ranging from ~ 10 pN (constructs that rupture through unzipping) to ~ 50 pN (constructs that rupture in shear). However, duplex constructs are irreversible. Recent efforts<sup>38</sup> have developed reversible construct designs by adding a loop that keeps two strands that form a single interaction

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pair tethered together in the open state, but these still rely on different devices for different forces, and devices are not easily exchanged since the constructs are generally directly attached to the glass surface. Furthermore, the limited stability of duplex DNA could limit their use in other biological environments. DO devices are more stable than duplexes<sup>39</sup> providing an advantage for some applications, and the ability to integrate multiple force-sensitive interactions in DO devices provides modularity to tune force response without the need for redesign of the primary structure. Prior work<sup>29</sup> has demonstrated the inclusion of multiple hairpins in a DO device allows for tuning the rupture response over the range of  $\sim 8-19$  pN, but these prior DO designs are not adjustable in real-time. DO devices provide the potential to be modified and tuned without the need for redesign of the primary structure and to be modified in realtime<sup>40</sup>. Here we take a distinct design approach where a base device is folded and then one or more ssDNA molecules are added to introduce one or more force-sensitive interactions after folding, or even after initial testing, to control the force-response of the device.

We focus on a DO nanodevice, the NanoDyn (ND), which has been previously shown to be sensitive to compressive depletion forces<sup>30</sup>. Hudoba et al. introduced the ND as a sensitive reporter of compressive depletion forces due to local molecular crowding on the order of 100 femtoNewtons (fN) and with a lower limit of force detection of 40 fN. Here, we build on that research and demonstrate the utility of the ND not only as a highly sensitive reporter of compressive depletion forces, but also as a robust, dynamic device capable of responding to tensile forces ranging from a few picoNewtons up to tens of picoNewtons (pN) where device design parameters allow tunable control of the force response.

Taking advantage of the modular nature of the ND, we show that an individual single stranded DNA molecule, which we refer to as a zipper strand, can be modified to set the force response and be incorporated after folding and purifying the ND. This allows for rapid and efficient tuning of the device and eschews the need to fold and purify a separate structure for different force applications. We investigated its response to tensile forces and determined that it can be tuned to be sensitive to a range of forces through the adjustment of 1 to 3 zipper strands. We show that the ND detection force can be adjusted between 5 and 13 pN by changing a single zipper strand within the device. We then demonstrate that by incorporating multiple zippers in parallel, the ND responds at forces of about 30 pN with the potential of even higher force induced opening. We find that more stable interactions (opening forces  $\gtrsim 10$  pN) lead to a higher reclosure probability. Finally, we show that the force response range of the ND can be adjusted in real time by iteratively incorporating DNA zippers in situ. This study lays the groundwork for a modular and versatile force responding probe that has the potential to be used in complex biological systems where traditional force spectroscopy techniques are challenging or impractical to implement.

#### Results

#### A DNA origami (DO) sensor design for a modular and tunable force response

As previously reported<sup>30</sup>, the ND is prepared by scaffolded DNA origami<sup>40-42</sup> where it consists of two origami bundles in a honeycomb lattice linked by six parallel 116 nucleotide (nt) single strand (ss) connections that we refer to as loops. Each loop is configured as either a "force-responding" loop or a "hinged" loop with the addition of ssDNA molecule(s) (Fig. 1a, Supplementary Fig. S1). A force-responding loop contains a zipper strand DNA oligonucleotide with 3 distinct regions, which allows the ND to transition between open and closed states. (i) The "anchor" region binds to one side of a given loop with 30 complementary bases so that this region remains base paired in both the open and closed states. (ii) The "zipper" region binds the opposite side of the loop such that the two origami bundles are constrained in the closed state. The length of the zipper region is varied between 11 and 21 nt to influence the opening force of the ND. (iii) The "linker" region is a 5 nt poly-T sequence linking the anchor and zipper regions, that helps reduce steric clash within the closed state of the ND. The hinged loops contain two separate 46nt DNA molecules referred to as "blocking strands" that form two dsDNA regions that are separated by 14 nt of ssDNA between them and leave a 5 nt ssDNA spacer adjacent to the ND barrels. This reduces the impact of secondary structure and entropic elasticity within the hinge loops on its force response. The modularity of the ND allows for the zipper strands to be incorporated after folding and purification of the base structure of the ND in a secondary reheating and slow annealing cycle (see "Methods" for details). This allows a single preparation of the ND base structure to be used for multiple ND force-response configurations. The ND folding, purification and zipper incorporation were verified via Agarose Gel Electrophoresis (AGE) (Supplementary Figs. S2, S3) and Transmission Electron Microscopy (TEM) (Fig. 1a, Supplementary Fig. S4).

To investigate the force response of the ND, we used a Magnetic Tweezers (MT) approach to carry out repeated force-extension measurements. Using a lab-built MT on an inverted Olympus IX-70 microscope base, we repeatedly actuated single NDs between the closed and open states by serially increasing and lowering the applied force. Each ND was attached to a 2.8 µm streptavidin coated superparamagnetic bead by 4 biotinylated dsDNA extensions at the top end of the ND (Fig. 1b). The opposite end of the ND was anchored to a  $\sim 1 \, \mu m$ dsDNA tether via base pairing between two complementary 30 nt ssDNA overhangs. The opposite end of the tether contained a 60 nt ssDNA overhang that anchored it to a glass slide by annealing to a complementary 60 nt ssDNA oligonucleotide that was covalently bound to the slide surface via click chemistry (see "Methods"). If all six loops of the device are either blocked (open) or closed by long zippers the devices show smooth force extension curves (Supplementary Fig. S5). However, for devices with between one and three force-responding loops, an abrupt increase in the extension of the ND by tens of nanometers occurred as the force was steadily increased (Supplementary Fig. S6). This indicated an opening event, where the zipper region within the force-responding loop released from the loop, leading to separation of the two barrel components. This gap size between the open and closed ND agrees with previously reported opening distances<sup>30</sup>. To allow the ND to reclose, the force was decreased to ~0.5 pN, which allowed the zipper region to rebind within the force-responding loop and close the ND. This also confirmed that the anchor end remained bound to the loop through any prior opening events.



**Figure 1.** DNA origami NanoDyn schematic and experimental design. (**a**) Schematic drawing of a DNA origami NanoDyn (ND) which consists of 2 honeycomb lattice barrels held together by 6, 116 nt ssDNA crossover strands called loops. Each loop can be folded into a "force-responding" loop or a "hinged" loop through annealing of a zipper strand or blocking strands, respectively. The force-responding loop can be in an open or closed state. TEM images provide visualization of these two states. (**b**) The ND is attached to a 2.8 µm superparamagnetic bead through 4 biotin-streptavidin linkages. The opposite end of the ND is annealed to a dsDNA tether, which itself is annealed to an oligonucleotide covalently bonded to a microscope slide via click chemistry. Repeated actuation of the ND is achieved by repeatedly increasing and then decreasing the force using a magnetic tweezers system. A sudden increase in length during the force loading step is indicative of an opening event.

Repeated extension-retraction cycles of multiple NDs resulted in a distribution of opening forces, which allowed identification of the median opening force for a given design.

### A single force-responding loop can tune the NanoDyn force sensitivity

As a two-state device the ND only reports if the applied force is above or below its force threshold. To investigate the tunability of this force threshold, we first focused on a ND with a single force-responding loop and varied the zipper region length within this single loop. A schematic of the design is shown in Fig. 2a, where a single loop (loop 6) was held closed by a zipper strand. We use the nomenclature 'L#' to indicate the loop being referenced, and '#nt' to indicate the length of the zipper region. For example: a 13nt zipper region in loop 6 is referenced as L6-13nt. The remaining 5 loops (L1-L5) were folded as hinged loops (Fig. 2a).

We used the MT to apply repeated force-extension measurements of NDs with a range of zipper region lengths in loop 6. We found that zipper region lengths of less than 13nt in L6 rarely closed. Therefore, we investigated NDs with 5 separate zipper region lengths in L6 of 13nt and longer: L6-13nt (blue), L6-15nt (green), L6-17nt (yellow), L6-19nt (orange), and L6-21nt (red). As the force was increased, the end-to-end extension of the DNA handles with the ND increased continuously with an abrupt extension that was due to the ND opening. Representative force-extension data for L6-13nt and L6-21nt are shown in Fig. 2b, while representative data for L6-15nt, L6-17nt, and L6-19nt are shown in Supplementary Fig. S7. Because the ND is directly attached to the bead, we develop a worm-like chain plus torsional spring (TorWLC) model to account for the torque applied to the ND as a result of randomized placement of the ND on the bead surface and the alignment of the magnetic moment of the bead with the externally applied magnetic field (see "Methods" and Supplementary Methods for details). The force response of the ND plus DNA handles with the rupture removed was fit to the model (TorWLC<sub>closed</sub>, Supplementary Fig. S8), and the post-rupture force response was derived from this fit by increasing the length of the opened ND (TorWLC<sub>open</sub>). This model does not explicitly describe the opening of the device itself but solely serves to verify single tether attachment and deduce the origin of the distance scale by describing the force-extension behavior of the entire assembly outside of the actual opening event.

We carried out multiple force-extension measurements with more than 10 molecules of each ND configuration, which resulted in the observation of more than 100 opening events for each configuration (Supplementary Table S1). We plotted the cumulative probability (Fig. 2c) and determined both the median force (Fig. 2d) and the opening distance (Supplementary Fig. S6, Supplementary Table S2) of each opening event. We found that the increase in the zipper region length in steps of two nucleotides from 13 to 21 nt correlated with an increase in the median opening force of 6.6 pN, 8.0 pN, 9.2 pN, 12.6 pN, and 13.0 pN, respectively (Supplementary Table S1). The standard deviation (SD) for each respective configuration was 1.7 pN, 1.2 pN, 2.8 pN, 2.1 pN, and 1.4 pN (Supplementary Table S1). The median, mean, standard error of the mean (SEM), and interquartile values for these opening force and distance measurements are provided in Supplementary Tables S1 and S2, respectively. The observation that most of the opening transitions occurred at or below 15 pN is consistent with previous



**Figure 2.** Zipper region length within a single force-responding loop modulates the opening force. (a) Schematic design for ND with one force-responding loop at L6 and five hinged loops at L1-L5. (b) Representative force-extension curves for L6-13nt (blue) and L6-21nt (red). Data was fit to a Torsional spring + worm-like chain model (TorWLC) for both the closed (TorWLC<sub>closed</sub>) and open (TorWLC<sub>open</sub>) states. (c) Cumulative probability distribution of the opening forces for multiple pulls across multiple devices. (d) Violin plots of the opening force distributions. The white dot is the median opening force value, which is indicated above or below each respective distribution and the black bar indicates the first quartile above and below the median. The median, mean, SD, SEM, and interquartile values for each ND version can be found in Supplementary Table S1.

studies that investigated the forces for unzipping DNA hairpin structures, which show that the force required to unzip DNA converges around 15 to 20  $pN^{29,43,44}$  depending on the buffer ionic conditions<sup>45–47</sup>. Overall, these results indicated that length variation of the zipper region within a single loop can finely tune the opening force between 6 and 13 pN, where the lower bound is set by the closure probability and the upper bound is the unzipping force of an infinite DNA duplex.

#### Parallel force-responding loops increase the force detect by ND opening

Given the limited range of opening forces the ND can detect with a single force-responding loop, we investigated the impact of including multiple force-responding loops arranged *in parallel* by the ND geometry (Fig. 3a). For the second loop, we focused on loop L3 because it is positioned opposite to loop L6, while for the third loop, we used loop L1 that is adjacent to loop L6. As the loops are different regions of the bacteriophage scaffold, they have different sequences. To help ensure similar opening force medians, we chose zipper region lengths, L1-13nt and L3-11nt so that their melting temperatures ( $T_m$ ) were similar (Supplementary Table S3) to L6-13nt. We first prepared the two additional ND versions with a single force-responding loop at L1-13nt or L3-11nt as controls for studies of NDs with multiple force-responding loops. We carried out force-extension measurements with the MT (Fig. 3b and Supplementary Fig. S7), plotted the cumulative probability (Fig. 3c), and found that the L1-13nt and L3-11nt NDs opened at respective median forces of 8.5pN and 5.5pN (Fig. 3d) with respective SDs of 2.1 pN and 1.2 pN (Supplementary Table S1). The opening force statistics are comparable to the opening forces measured for the L6-13nt ND.

We then prepared a ND with both L6-13nt and L3-11nt, which we refer to as L6+L3, and investigated the force required to open this two-zipper ND (Fig. 3c,d, Supplementary Fig. S7, Supplementary Table S1). We carried out force measurements on 23 ND devices and detected a total of 205 opening events (Supplementary Table S1). We observed a single step with an opening distance that was nearly identical to the opening of the ND with either L6-13nt or L3-11nt (Supplementary Fig. S6, Supplementary Table S2). This is likely because once the first loop completely ruptured, the force was significantly above the force that the second (potentially partially) closed loop can support, so it ruptured faster than can be detected by the MT instrument.

For each individual device, there was the possibility that only one of the two zipper strands was incorporated, which would result in a ND that contained only one force-responding loop. To address this possibility, we verified that each two-zipper ND contained both zipper strands by using log-likelihood analysis (see "Methods" for details). Briefly, by using the two force distributions of a ND with 1 force-responding loop as the reference model, we determined if any of the force distributions of an individual ND fit better to a one-zipper distribution. Any such devices (5 out of 28 total molecules) were not included in the two-zipper ND analysis. This implies that



**Figure 3.** Multiple force-responding loops increase the ND opening force. (**a**) Schematic design for force-responding loops with (i) one force-responding loop at loop 6 (blue), 3 (purple), and 1 (grey); (ii) two force-responding loops at loops 6 and 3 (pink); and (iii) three force-responding loops at loops 6, 3, and 1 (dark red). (**b**) Representative force-extension curves for L3-11nt (purple) and L6-13nt + L3-11nt + L1-13nt (dark red—abbreviated L6+L3+L1). Data was fit to the TorWLC model. (**c**) Cumulative probability distributions of the opening forces for the ND devices. (**d**) Violin plots of the opening force distribution for each ND device. The white dot indicates the median opening force, which is shown above or below each respective distribution and the black bar indicates the first quartile above and below the median. The median, mean, SD, SEM, and interquartile values for each ND version can be found in Supplementary Table S1.

82% of the NDs contained 2 force-responding loops. A histogram comparing the pre and post log-likelihood analysis data is shown in Supplementary Fig. S9.

After removal of the devices with a single force-responding loop, we plotted the cumulative probability of the two-zipper ND opening as a function of force (Fig. 3c). We found that the device opened with a single step at a median force of 11.1pN with a SD of 2.8 pN (Fig. 3d and Supplementary Table S1). For comparison, we plotted the normalized sum of the cumulative probabilities of the ND's with the single force-responding loops with either L6-13nt or L3-11nt (Supplementary Fig. S10, "Methods"). We found that this inferred cumulative probability was within 9% of the two-zipper ND that contained both L6-13nt and L3-11nt. Overall, these results indicate that using the ND geometry to orient two force-responding loops in parallel results in an additive increase in the opening force of the ND, which significantly expands the force-responding range of the ND. This is somewhat different from a previous study<sup>29</sup> that used a DO to orient DNA hairpins in parallel. They found that the force to open the hairpins increased but that the increase in force was less than additive. This suggests the ND allows for a more even distribution of forces between zippers.

To build off the results from combining two force-responding loops in the ND, we included a third zipper strand to introduce a third force-responding loop, L1-13nt. We carried out force-extension measurements of the three-zipper ND (Fig. 3b), which we refer to as L6 + L3 + L1. We observed a single opening step, as was observed with L6 + L3. This is again consistent with the idea that the three force-responding loops in parallel support a high enough force that once one of the zipper regions ruptures, the force on the two remaining loops is high enough so they rupture faster than the time resolution of the MT measurement.

We used log-likelihood analysis (see "Methods" for details) to verify if a L6 + L3 + L1 ND contained three force-responding loops, as was done with the L6 + L3 ND. We compared the force opening distributions of individual L6 + L3 + L1 ND to the distributions of each ND with a single force-responding loop, and the corrected distribution of the L6 + L3 ND. We found that 3 out of the 14 devices studied had a distribution that more likely contained two force-responding loops. Those 3 devices were therefore removed from further analysis. Hence, 11 out of 14 L6 + L3 + L1 NDs, or 79%, contained all three force-responding loops. A histogram comparing the pre and post log-likelihood analysis data is shown in Supplementary Fig. S9. There remains a small fraction of opening events that appear to align better to the L6 + L3 distribution than the L6 + L3 + L1 distribution. This is likely because when ND reclosed, not all the force-responding loops reclosed each time. Importantly, this was a small fraction of the total opening events.

After correcting for NDs that contained less than three force-responding loops, we determined the cumulative probability for the L6 + L3 + L1 ND to open (Fig. 3c) and found that the median opening force was increased to 25.9 pN with a SD of 6.1 pN. We compared these results to the opening force distribution that was inferred from summing the cumulative probabilities of each of the three individual zippers (Supplementary Fig. S10), which implied a median opening force of 15.4 pN with a standard deviation of 4.3 pN. This inferred median force was

about 40% lower than the measured value, suggesting that additional interactions were introduced into the device by the addition of the third force-responding loop in L1. Interestingly, the median step size for L6 + L3 + L1 ND (38.8 nm) was significantly larger (Supplementary Fig. S6, Supplementary Table S2) than all the other ND versions we studied including the L6 + L3 ND (31.3 nm). Since the length of the ND in the open state is given by the extension of the blocked loops and thus will be similar across all configurations, this increase in step size of the L6 + L3 + L1 ND was likely due to its length in the closed state being shorter than the other ND versions including the L6 + L3 ND. This could be due to base stacking interactions between the two barrels of the ND that is known to occur between DNA origami devices<sup>27,48,49</sup> even though the end connections of adjacent dsDNA helices contain ssDNA loops that suppress base stacking interactions. Furthermore, we saw a slight correlation between the opening force and opening distance for L6 + L3 and L6 + L3 + L1 that was not present in the ND's with only one force-responding loop (Supplementary Fig. S11). This is consistent with the idea that additional interactions such as base stacking shorten the length of the closed ND and result in a higher opening force (correlations between opening distance and rupture force due to the force-dependent extension of the blocking strands and due to the force-dependent tilt of the NanoDyn would be expected to be more pronounced at the lower forces where both effects are less saturated). Overall, these results demonstrate the versatility of the ND, where integrating multiple force-responding loops in parallel allows the ND to detect forces higher than the inherent limit of single dsDNA unzipping forces, expanding the versatility of this nanoscale device for force-responding applications.

#### Increased interaction stability improves the closure efficiency

For the ND to function as a reversible force sensor that measures repeated application of an external force, it needs to reclose efficiently following release of the applied force. To investigate this, we determined the fraction of times each ND closed (and subsequently reopened) following a reduction in the applied force to 0.5 pN (Fig. 4a). We typically did not directly observe a closing event as the force was reduced. This is likely because the closing events usually occurred at forces where the bead height fluctuations were comparable to the closing step size of ~ 30 nm, making the closing step difficult to detect (Supplementary Fig. S12). Furthermore, these observations imply that observing multiple opening and closing events at a constant force is not feasible with these ND devices. So instead, we determined the fraction of force-extension experiments with a subsequent opening event after the force was reduced to 0.5 pN. We found that the opening efficiency of NDs with one force-responding loop improved as the zipper region length (and force to open) increased (Fig. 4 and Supplementary Fig. S13). For 11 nt and 13 nt zipper regions, the closing efficiency was less than 50% (Fig. 4, Supplementary Table S4). An



**Figure 4.** Effect of zipper region length and multiple force-responding loops on ND closure and subsequent re-opening. (a) Violin plots of the fraction of extensions with an opening event for each individual ND. The triangles indicate the position in the distribution of the devices shown in (**b**–**d**). The white dot indicates the median opening fraction, which is indicated above or below each respective distribution. The black bar indicates the first quartile above and below the median. The median, mean, SD, SEM, and interquartile values for each ND version can be found in Supplementary Table S4. (**b**–**d**) Examples of repeated force-extension curves of single (**b**) L1-13nt, (**c**) L6-13nt, and (**d**) L6+L3+L1 ND devices. Each plot contains 10 consecutive extensions with arrows indicating opening events. The corresponding retraction cycles can be found in Supplementary Fig. S12.

increase in the zipper region length in L6 to 15 nt, 17 nt, 19 nt, and 21 nt resulted in closing efficiencies of 57%, 61%, 99%, and 82%, respectively (Supplementary Table S4). This generally implies that increasing the zipper length improved closing efficiency. To rule out the possibility of zipper strand loss over many force-extension cycles of the ND contributing to a lower closing efficiency, we plotted the fraction of devices that re-opened as a function of the cycle number (Supplementary Fig. S14). We found that over 8 force-extension cycles, a significant drop in the fraction of devices that re-opened was not observed, indicating that it is unlikely that zipper strands were lost over the course of the experiment, again confirming that the anchor region effectively stayed bound through successive opening events. In the case of L6-21nt, the closing efficiency decreased relative to L6-19nt. One possible explanation is that the 5-prime end of the zipper region in L6-21 starts with several A's that could transiently bind to the poly-T region of the zipper strand and interfere with proper rebinding. We determined the opening fraction of L6+L3 and L6+L3+L1 NDs to assess the impact of multiple force-responding loops (Fig. 4). Interestingly, we found that including multiple force-responding loops reliably increased the closing efficiency to 91% and 94% for the L6+L3 and L6+L3+L1 NDs, respectively. These results indicate that increased stability by either an increase in zipper region length, or inclusion of multiple force-responding loops improves closing efficiency.

#### Real-time incorporation of DNA zippers into a single ND to actively control force response

We have shown that the force the ND detects relies on the type and number of force-responding loops it contains. The zipper strands for the preceding experiments were incorporated after folding of the base structure of the device, which allowed the opening force to be defined through the addition of just a few DNA oligonucleotides prior to use in an experiment. To investigate the potential for real-time modulation of the opening force, we chose an ND design that allowed for successive integration of multiple zipper strands; one to define the initial opening force and another to adjust the opening force of the same single device (Fig. 5a).

To demonstrate this approach, we focused on the L6-13nt and L3-11nt zippers, which had median opening forces of 6.6 pN (SD 1.7 pN) and 5.5 pN (SD 1.2 pN), respectively. Starting with an unconstrained ND (omitting blocking strands in L6 and L3), we found that force-extensions never resulted in well-defined opening events, as expected (Fig. 5b). We then introduced 800 nM of the L6-13nt zipper strand into the flow chamber and incubated for 20 min with a 1 pN applied force on the ND (see "Methods" for details). After flowing out the excess unbound zipper strands, force-extension measurements revealed a single well-defined step in the expected force range (Fig. 5c). The opening force was within the force distribution of the L6-13nt ND presented above.



**Figure 5.** Iterative introduction of multiple force-responding loops into a single ND. (**a**) Diagram of the iterative introduction of one and then two force-responding loops into the ND. (**b**) Force-extension of the ND prior to incorporation of a force-responding loop never results in an opening event. (**c**) Following incubation with a single zipper strand that forms the L6-13nt force-responding loop, a well-defined step is observed upon force-extension. (**d**) Following incubation with a second zipper strand that forms the L3-11nt force-responding loop, an additional increase in the opening force is observed.

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To adjust the force-responding range, we then incubated the ND with a second zipper strand, L3-11nt, 800 nM for 20 min and then flowed out any excess zipper strand. Force-extension experiments revealed that the addition of this second zipper strand resulted in an increase in the opening forces to ~9–13 pN, which is consistent with previous 2-zipper ND measurements (Fig. 5d). We found that the first and second zipper strands incorporated with an efficiency of 25% and 23%, respectively. Future work beyond the scope of this study will be necessary to achieve more efficient incorporation of multiple force-responding loops. However, as a proof-of-concept, these results reveal an important potential avenue for the ND, which is the ability to incorporate zipper strands into the ND in situ, allowing for the force-responding to be initiated and then adjusted within a single device during an experiment.

#### Discussion

In this work, we demonstrated that the DNA origami ND can function as a modular nanodevice that can be tuned to detect a range of tensile forces. The modular design allowed the base-structure to be folded, purified, and stored without force-responding loops. Then, immediately before use, the force detection range was customized by incorporating one or more zipper stands. We demonstrated that varying the zipper region length within a single force-responding loop modulated the opening force over a range of threefold up to a maximum unzipping force of about 15 pN<sup>43,44</sup>. We then showed that including multiple force-responding loops in parallel enabled a wider range of opening forces up to 26 pN with only 3 relatively weak zippers, which exceeds the inherent unzipping force of 15 pN. We found that the interaction stability affected the reversibility of ND, with median opening force cycling. However, devices with a median opening force above 10 pN repeatedly reclosed, which confirms their utility in detecting repeated force applications. Finally, we showed that single DNA zipper strands can be iteratively incorporated into the ND during a force measurement. This opens the possibility for tuning the force-responding range of single NDs in real time during a measurement.

This work expands the utility of using nanoscale force sensors as a complementary approach to existing force spectroscopy techniques. The ND has the potential to probe a wide range of forces in constrained environments where it can be difficult to implement other force spectroscopy techniques<sup>4,6-12</sup>. We previously showed that the ND can operate in crowded environments and detect compressive depletion force in the range of 0.05 to 1 pN<sup>30</sup>. Here in this work, we demonstrated the same ND base structure can also be used to detect tensile forces from 6 to at least 26 pN. The ability for the ND to operate in different modes for detecting both compressive and tensile forces with order of magnitude different force ranges indicates its high versatility for a DNA origami device<sup>23,25,26,29-31,50,51</sup>. In comparison to the previous study presented in Dutta et al.<sup>29</sup>, our results indicate the ND provides a wider dynamic range of force sensing. This is consistent with the idea that integrating the forceresponding loops between the two-barrel structures allows for a more balanced distribution of the force on these force-responding loops. There is the potential for further versatility of the ND since up to six force-responding loops could be included in the ND, each with independent nucleotide sequences to which DNA zippers can be incorporated independently and reproducibly. It will be important to ensure efficient incorporation of all zipper stands. While we achieved full incorporation of 3 zippers within about 80% of NDs, further optimization will be important as additional force-responding loops are used. However, the number of zippers could be directly detected with single molecule fluorescence and photobleaching, which could alleviate the need for further optimization.

Assuming an opening force of 10 pN of force per force-responding loop, using six force-responding loops should result in an opening force of more than 60pN of force. These large forces do occur in biological systems including the forces on phage genomes during viral packaging<sup>52,53</sup> and the forces on mitotic chromosomes during mitosis<sup>54</sup>. However, measurements of these high forces will require covalent attachments or multiple non-covalent attachments to prevent failure of the attachment before device opening and force detection<sup>13,55</sup>.

The overall length of the ND at 100 nm in length is advantageous for constrained environments. However, for experiments requiring smaller devices, the overall length of the ND could be reduced by designing shorter barrels, while retaining the loop regions. The shortened length could be accomplished with the same DNA scaffold by increasing the width, or with a shorter DNA scaffold. In addition to our current method of monitoring relative length change with magnetic tweezers, a fluorophore pair that undergoes Förster Resonance Energy Transfer (FRET) can be incorporated into the ND with 2 fluorophore labeled oligos, as reported in Hudoba et al.<sup>30</sup>, where high FRET reports a closed state and low FRET reports the open state. This will allow detection of a force range in environments where attaching a force handle is not possible.

In the broader context of applications for the ND, there is significant potential for investigating cell-cell and cell-surface interactions based on previous studies. ssDNA hairpins<sup>36,37</sup> and DNA origami platforms<sup>29</sup> have been successfully implemented to investigate intercellular forces. The ND could be used similarly where the modularity of the ND could complement these previously published elegant studies by enabling a wider range of forcesensing. Furthermore, iterative zipper incorporation would allow the force sensor to be tuned in conjunction with changes in the extracellular environment that cause the cells to adapt by changing their cell-surface interactions. Future studies will be needed to investigate these potential applications of this versatile nanoscale device.

#### Methods

#### Preparation of the DNA origami NanoDyn (ND)

The ND design, as previously described in Hudoba et al.<sup>30</sup>, consists of 2 DNA bundles connected by 6 crossover strands. One bundle contains 24 dsDNA helices bundled in a honeycomb pattern and has a 30 base ssDNA protruding from the outer end to facilitate attachment to a dsDNA tether. The second bundle contains 18 dsDNA helices arranged in the same fashion as the first bundle but with the central 6 helices omitted. This bundle has 4 biotinylated dsDNA overhangs arranged around the periphery of the outside end (opposite the tethering end) which facilitates attachment to a streptavidin labeled bead. The overall dimension of the ND is ~ 100 nm × 15 nm as measured by Transmission Electron Microscopy (TEM). The caDNAno design for this structure is provided in Supplementary Fig. S1.

The backbone of the ND consists of an 8064 base scaffold from a modified M13mp18 bacteriophage sequence produced in Castro lab<sup>40</sup>. ~ 170 oligonucleotide staples (Integrated DNA Technologies) were combined with the scaffold to fold each version of the device. The exact number of staples is dependent on the version of ND since the number of loops into which zippers are later incorporated will also affect the blocking strands that will be included in the main folding of the ND. Oligo sequences for the base structure can be found in Supplementary Table S5, specific zipper sequences in Supplementary Table S3, and loop sequences (scaffold crossover points) in Supplementary Table S6. The folding was carried out as described in previous folding protocols<sup>40</sup>. In brief, 100 nM scaffold is combined with 10× staple strands in a folding buffer (5 mM Tris, 5 mM NaCl, 1 mM EDTA, 18 mM MgCl<sub>2</sub>, pH 8). The reaction is then subjected to a thermal annealing ramp starting at 65 °C to disrupt non-specific base pairing interactions and then slowly cooled in incremented temperature steps until 4 °C. The detailed thermal ramp is contained in Supplementary Table S7. The folding was confirmed by agarose gel electrophoresis (AGE) (Supplementary Fig. S2) as previously described<sup>40</sup>. The resulting samples were polyethylene glycol (PEG) purified to remove excess staple strands (Supplementary Fig. S2). The PEG purification is based on existing protocols<sup>56</sup>. Briefly, the folded ND sample was combined at a 1:1 volumetric ratio with 15% PEG MW 8000 (Sigma Aldrich) suspended in folding buffer with added 500 mM NaCl. The sample was then pelleted by centrifugation at 16,000g for 30 min at room temperature. The supernatant was removed and the pellet resuspended in folding buffer with 10% glycerol, aliquoted, flash frozen, and stored at -80 °C<sup>57</sup>.

After folding the base structure of the ND, individual zippers are subsequently incorporated by incubating 4 nM ND with a 10× concentration of the desired zipper(s) for 1 h at 45 °C then cooling by 1 °C/min until it reaches 10 °C. This reaction takes place in the same folding buffer described above. In the initial folding, it is recommended to omit blocking strands (strands meant to make the ss scaffold loops double stranded to block unwanted non-specific interactions) only from loops that are intended to be used in subsequent zipper incorporation. The zipper incorporation was confirmed by AGE (Supplementary Fig. S3) and TEM (Supplementary Fig. S4).

#### Preparation of a dsDNA tether handle

Tethers were prepared by restriction enzyme digestion of pUC19 with BsaI (New England Biolabs: R0535) in  $1 \times$  CutSmart buffer. 1–3 units of enzyme per 1 µg of DNA is usually sufficient to digest the plasmid without over-digestion. The digestion takes place at 37 °C for 1 h followed by a heat shock at 65 °C for 20 min to inactivate the enzyme. The digestion is verified by AGE (Supplementary Fig. S15). The gel conditions are as follows and are used for all gels in the tether preparation: 0.7% agarose gel run in 0.5 xTAE at 225 V and post stained with ethidium bromide for UV visualization. The linearized plasmid was subsequently ligated to oligo pairs to create long ssDNA overhangs (Sigma-Aldrich) on either end. One DNA end has a 3' 30 nt ss overhang to facilitate attachment to the NanoDyn and the other end has a 3' 60 base ss overhang to attach to an oligonucleotide covalently bound to the microscope slide. Prior to ligation, the 5' end of each oligonucleotide to be ligated was phosphorylated using T4 Polynucleotide Kinase (T4PNK) (New England Biolabs: M0201) at 1 U/25 pmol ends incubated at 37 °C for 90 min followed by a 65 °C heat shock for 20 min. This reaction was performed in 1×T4 ligase buffer rather than T4PNK buffer since they would be subsequently ligated to the linearized plasmid. Each of the 2 oligo pairs were separately annealed at an equal molar ratio at room temperature for 15 min in 0.5× TE with 50 mM NaCl (Sigma-Aldrich). The 2 pairs of ends were then ligated simultaneously to the linearized plasmid using T4 DNA Ligase (New England Biolabs:M0202) at 2 U/pmol DNA ends in 1×T4 ligase buffer (provided with enzyme). 100-fold excess ends were used during ligation to prevent recyclization or oligomerization of the linearized plasmid. Ligation was verified by AGE (Supplementary Fig. S15). Following the ligation, a phenol chloroform extraction was performed to remove bovine serum albumin (BSA) contained within the digestion reaction buffer. The tether was purified away from excess ends via high pressure liquid chromatography (HPLC) using a Gen-Pak column (Waters: WAT015490). The desired sample fractions were identified by AGE. Those fractions were combined, and the final tether sample was then buffer exchanged and concentrated into  $0.5\times \mathrm{TE}$ using a 30 kDa centrifugal filter (Millipore-Amicon Ultra).

#### Preparation of single molecule experiment slides

The flow cell preparation is an adaptation of the methods used in Luo et al.<sup>58</sup> and Chandradoss et al.<sup>59</sup>. First, holes are sandblasted into either end of a coverslip creating an entry and exit port for the sample. This 'top' coverslip is then thoroughly rinsed with Milli-Q water to remove excess sand. The top and an equal number of 'bottom' coverslips are then sonicated in isopropyl alcohol for 20 min. Both sets of coverslips are then thoroughly rinsed in Milli-Q water. The bottom coverslips are dried in an 80 °C oven while the top coverslips are submerged in a 1% Hellmanex III solution (Sigma Z805939), brought to a boil in the microwave, sonicated for 20 min, thoroughly rinsed in Milli-Q water, then placed in the 80 °C oven to dry. When the bottom coverslips are dry, they are plasma cleaned (Electron Microscopy Sciences: K100X) for 4 min at 25 mA for surface activation. Immediately following the plasma cleaning, the bottom coverslips are incubated in a 3/100 mixture of 3-aminopropyltriethoxysilane (MP Biomedicals: 02154766) and acetone (Sigma 650501) for 30 min in a nitrogen gas filled desiccator. Following silanization, the coverslips are sonicated in fresh acetone for 5 min, thoroughly rinsed with Milli-Q water, and placed in the 80 °C oven to dry. The top and bottom coverslips are sandwiched around parafilm with a channel cut into it. The sandwich is heated to ~75–80 °C to melt the parafilm, adhering the coverslips and creating the flow channel.

The following steps describe a click-chemistry procedure to covalently attach a DNA oligonucleotide to the slide. The reaction mixture consists of a 10% mPEG-SVA (Laysan Bio: MPEG-SVA-5000) solution in 0.1 M potassium tetraborate pH 8.1. To that, DBCO-PEG4-NHS ester (Conju-Probe: CP-2028) at 400  $\mu$ M resuspended in anhydrous DMSO (ThermoFisher Scientific: D12345) is combined at a 40:1 ratio with a 10  $\mu$ M azide modified oligo (Sigma-Aldrich) for a final concentration of 50 nM DNA in the reaction mixture. The reaction mixture is injected into the flow cell and incubated for 1 h at 23 °C. It is subsequently rinsed from the flowcell with Milli-Q water and the same reaction and incubation is performed a second time using fresh reagents. After the second incubation, the flow cell is rinsed thoroughly with Milli-Q water and allowed to dry over 18–24 h at 23 °C in a nitrogen storage box. After the flowcells are dry, they are individually vacuum sealed and frozen at -20 °C until the day of the experiment.

On the day of the experiment, one flowcell is removed and brought to room temperature before removal from the packaging. The surface is passivated with Blocking Reagent (Roche: 11096176001) resuspended in 0.5 xTE and allowed to incubate for 20 min before rinsing with the experiment buffer (0.5 xTE with 100 mM NaCl, 10 mM MgCl<sub>2</sub>, and 0.5% Tween20). Separately, the ND is annealed to the tether by incubating at 45 °C for 1 h and then cooled at 1 °C/min. The annealing is checked using AGE (Supplementary Fig. S16). The ND + tether construct is then incubated with streptavidin coated superparamagnetic particles (prerinsed in experiment buffer) (Thermo-Scientific: Dynabeads M-280) for 15 min at room temperature. Pre-rinsed 4.5  $\mu$ m streptavidin labeled polystyrene particles (Spherotech SVP-40–5) are combined with 1  $\mu$ M biotinylated oligoes that are a reverse complement to the oligo covalently attached to the slide and incubated for 20 min at room temperature. Excess DNA is washed from the sample by repeated centrifugal pelleting, supernatant removal, and resuspension. These beads will adhere to the slide surface through many DNA's on the bead surface base pairing with the DNA on the slide surface and act as a fiduciary mark during the experiment. Just before injection onto the slide, the ND + tether + Dynabead construct is combined with the DNA coated fiduciary particles.

#### Single molecule force spectroscopy experiments on a magnetic tweezer system

Experiments were performed on a home-built magnetic tweezer system using an Olympus IX-70 inverted microscope body. Data collection for multiple samples is collected in parallel with an in-house written LabVIEW program. Prior to data collection, each bead position is individually calibrated so that relative position change of each bead can be measured. The force-extension is performed by moving a pair of permanent neodymium magnets close to the sample at a rate of 0.1 mm/s yielding loading rates of on the order of 0.15 pN/s at low (5 pN) forces and 0.85 pN/s at 20 pN. While the observed opening forces depend weakly on the rate of magnet movement, the rate of 0.1 mm/s was chosen as a compromise that limits the noise due to the device fluctuating at one force for a long time at low rates and minimizes artificially increasing the observed opening forces by driving the system further out of equilibrium (Supplementary Fig. S17). The magnets are then retracted at the same rate. This process can be repeated many times to collect multiple opening events from the same tethered sample.

The force exerted on each molecule is calculated using the equipartition theorem and the approximation of the bead-tether system as an inverted pendulum. These two concepts result in the equation:

$$F = \frac{k_b T L}{\langle x^2 \rangle} \tag{1}$$

where  $k_b$  is the Boltzmann constant, T is the temperature, L is end-to-end distance of the molecule, and x is the deviation of the bead from the central position due to Brownian fluctuations.

#### Modeling the NanoDyn as a torsional spring

To fit the experimental force extension curves, a model of the mechanical behavior of the ND + tether system was created. The model uses a worm-like chain for the tether, attached on one end to the surface and at the other end to a torsional spring, which represents the ND and its attachment to the bead. To avoid numerical artifacts due to the finite maximal extension of this model, we added a soft constraint on the physical length of the ND. We can then fit the force-extension curve predicted by the model to the experimental force-extension curves, by varying the three parameters of the model, namely the unknown offset in the distance measurements between the surface and the bead, the spring constant of the torsional spring, and the initial angle  $\Theta_0$  of the torsional spring with respect to the surface attachment point of the tether. A diagram of the system can be found in Supplementary Fig. S8. To account for the rupture, we first subtract the rupture from the high force regime data to give us a continuous data set to fit our low regime model to. We can then use the force point when the rupture occurred and the change in extension relative to the total length of the tether to algebraically calculate the newly acquired additional length of the system, adding that to the tether length to give us our high force regime fit. We note that the model does not predict the rupture itself but only describes the mechanical properties of the tether + device + bead assembly outside of the rupture event. A table of the fit parameters can be found in Supplementary Fig. S8. An extension of the model that introduces the additional post-rupture length of the flexible parts of the assembly and the effective length of the remaining fixed part of the ND as additional fit parameters can be fit to the data without removal of the rupture (Supplementary Fig. S18). The fitted effective lengths of the fixed part cover the entire range between 50 and 100 nm (Supplementary Fig. S18), which indicates that the connection between the two barrels of the ND is not fully flexible even in the open state (the length of the fixed part would be 50 nm, the length of the remaining barrel, if it was). Since the five parameter model does not visibly improve the quality of the fits over the three parameter model, we use the three parameter model in the main figures. A more detailed summary of the fitting algorithm can be found in the Supplementary Methods.

#### Flow incorporation of zippers into the ND in real-time

The setup of this experiment starts as previously described in the above methods. In the case of real-time zipper addition to the ND, a sample is prepared with the blocking strands omitted from the desired loops. To facilitate flow, the flowcell is attached via peristaltic pump tubing to a 1 mL syringe in a syringe pump (SyringePump.com). The pump is run at a rate of  $10 \,\mu$ L/min to draw sample through the ~  $30 \,\mu$ L flowcell sample chamber which is fed by a reservoir. To incorporate a zipper,  $200 \,\mu$ L of 800 nM zipper oligonucleotide is added to the reservoir and drawn through the sample chamber for 7 min with the magnets exerting a force of ~ 5 pN on the sample tethers, followed by a 20 min incubation at ~ 1 pN. During incubation, the reservoir is washed of remaining zipper by buffer exchanging 5 times with experiment buffer. Following the incubation, excess zipper is removed from the flowcell by drawing experiment buffer from the reservoir for 7 min with ~ 5 pN force applied to the sample.

#### Sample imaging by transmission electron microscopy (TEM)

For TEM imaging, structures were stained using a 1% Uranyl acetate solution and imaged on a FEI Tecnai G2 Spirit electron microscope. For each preparation,  $6-8 \mu l$  of 1 nM DNA origami sample was wicked onto a glow-discharge-cleaned copper grid (Electron Microscopy Sciences, Hatfield, PA) and incubated for  $5-10 \mu m$ . The sample solution was then removed carefully with Whatman #4 filter paper and the grids were immediately stained with two 6  $\mu l$  drops of 1% Uranyl acetate solution. Grids were dried for at least 10 min before imaging. TEM images were analyzed by ImageJ to try to quantify the length of the gap between the two barrels to determine if the L6 + L3 + L1 had a smaller gap at zero force than the other devices. However, the angular distribution prevented the quantification of the gap length (Supplementary Fig. S19).

## Log-likelihood removal of data from ND devices indicated as having incomplete zipper incorporation

We begin by using the 1-zipper distributions as reference models to determine if data from a single ND designed to have 2 zippers is "better fit" by the 1-zipper model. The 1-zipper distribution is assumed to be "correct" given the presence of the opening events. For each 2-zipper ND, the following calculation is performed:

#### $Log_{likelihood} = log(P(F_1)) + log(P(F_2)) + \dots + log(P(F_N))$ (2)

where P(F) is the normalized probability of an opening force, F, being in a given distribution. Here  $F_1$ ,  $F_2$ , ...,  $F_N$  are the N opening events observed for a given individual device. To account for 0 count bins in the probability distributions, we added + 1 pseudo-counts to each bin and renormalized the distributions accordingly. Not doing so results in undefined terms in the summation. This calculation is performed three times; once for each of the 1-zipper distributions (which are the 2 individual zippers contained in the 2-zipper ND) and once for the 2-zipper distribution. Smaller probabilities are more heavily weighted towards larger negative values, so whichever of the 3 calculated values results in the least negative number (closer to zero) is declared to be the distribution best fitting that list of forces. After testing all 2-zipper devices in this way, any of the 2-zipper devices indicated as being better fit by a 1-zipper distribution were removed from the aggregate data. We then repeated the same process for the 3-zipper ND data using the three 1-zipper distributions, the 'cleaned-up' 2-zipper distribution, and the 3-zipper distributions, for a total of 5 outputs. Again, any of the 3-zipper devices indicated as only having 1 or 2 zippers, were removed from the aggregate data. Histograms showing the data removal can be found in Supplementary Fig. S9.

## Estimate of the cumulative probability distribution for a ND with multiple force-responding loops

To estimate the expected cumulative probability distribution (CPD) for a ND containing multiple force-responding loops we added the CPD's of the individual force-responding loops. We assume that the force, F, is divided equally across all force-responding loops and that once one of the loops is opened, they all open due to the increase in force on the remaining zipper(s). This is supported by experimental data in that we do not see multiple smaller opening steps during force loading. For a ND with 2 force-responding loops, we calculate the cumulative probability that both zippers release at F, which is equivalent to the cumulative probability,  $P_c$ , of one zipper or the other breaking at F/2 given our assumptions. This results in the equation:

$$P_c(opening) = S\left(\frac{F}{2}\right) + R\left(\frac{F}{2}\right) - \left[S\left(\frac{F}{2}\right) \cdot R\left(\frac{F}{2}\right)\right]$$
(3)

where *S* and *R* are the CPD's for the two individual force-responding loops. We use a similar line of thinking for the ND with three force-responding loops where *F* is distributed evenly across three zippers. This results in:

$$P_{c}(opening) = S\left(\frac{F}{3}\right) + R\left(\frac{F}{3}\right) + T\left(\frac{F}{3}\right) - \left[S\left(\frac{F}{3}\right) \cdot R\left(\frac{F}{3}\right)\right] - \left[T\left(\frac{F}{3}\right) \cdot S\left(\frac{F}{3}\right)\right] - \left[T\left(\frac{F}{3}\right) \cdot R\left(\frac{F}{3}\right)\right] + \left[T\left(\frac{F}{3}\right) \cdot R\left(\frac{F}{3}\right) \cdot S\left(\frac{F}{3}\right)\right]$$
(4)

where *T* is the CPD for a third force-responding loop.

#### Data availability

The experimental data sets are either included in this submission, the supplemental information, or are available from the authors upon request.

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#### **Author contributions**

A.R. prepared the DO samples, DNA tethers, and flow cells, carried out force spectroscopy measurements, and data analysis; H.H. developed the TorWLC model; M.N. carried out the TEM imaging; P.B. provided support for the DO sample preparation; M.G.P, R.B., and C.E.C. supervised the study and helped with data interpretation; A.R. and M.G.P. wrote the manuscript; All authors edited the manuscript and approved the final version.

#### **Competing interests**

The authors declare no competing interests.

#### Additional information

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