DNA-Origami-Based Fluorescence Brightness Standards for Convenient and Fast Protein Counting in Live Cells

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Abstract

Fluorescence microscopy has been one of the most discovery-rich methods in biology. In the digital age, the discipline is becoming increasingly quantitative. Virtually all biological laboratories have access to fluorescence microscopes, but abilities to quantify biomolecule copy numbers are limited by the complexity and sophistication associated with current quantification methods. Here, we present DNA-origami-based fluorescence brightness standards for counting 5–300 copies of proteins in mammalian and bacterial cells, tagged with fluorescent proteins or organic dyes. Compared to conventional quantification techniques, our brightness standards are robust, straightforward to use, and compatible with nearly all fluorescence imaging applications, thereby providing a practical and versatile tool to quantify biomolecules *via* fluorescence microscopy.

Fluorescence microscopy is widely used in biology and biomedicine as a tool to visualize cell morphology and dynamics, in particular the biomolecules (e.g., DNA, RNA, protein) of interest that are labeled with a fluorophore. In modern life science studies, it is increasingly important to not only know the identities and locations of biomolecules, but also to measure their stoichiometry. However, methods to count biomolecules from fluorescence images usually lack versatility (e.g., they are limited to certain imaging modalities or dynamic range) and/or require sophisticated analysis. Stepwise photobleaching, while precise under finely tuned conditions, is limited to small (≤ 10) numbers of molecules.^{1–6} Quantitative immunoblot-calibrated internal fluorescence standards can quantify molecules of a wide range of copy numbers (from tens to thousands)⁷⁻¹⁰ but may suffer batch-to-batch variations that necessitate repeated, laborious calibrations. Super-resolution methods offer an increase in resolution and precision, but often at the expense of ease of use (i.e., specialized sample preparation and/or microscope) and speed (longer data acquisition and processing time).^{11–14} Therefore, there is a pressing need for a fast, universal technique that can be conveniently integrated into a wide array of existing imaging workflows to count biomolecules. A customizable external standard would fit this description: it would require little to no additional cloning, could be programmed to count virtually any fluorophore, and sustain long-term storage before being imaged in physiologically relevant conditions. As a precise, programmable, and robust self-assembly method, DNA origami^{15–18} is ideal for the creation of such a standard. DNA origami has been used to engineer standards for fluorescence microscopy, but existing quantification methods either rely on super-resolution techniques or only demonstrate applications for counting organic dyes.^{12,19} Here, we present DNA-origami-based brightness standards that incorporate 5-200 organic dyes and fluorescent proteins and use them for guantitative fluorescence microscopy on bacterial and mammalian cells. Our fluorescent standards offer a fast and precise solution to counting biomolecules in different cell types, with commonly used microscopes and fluorophores, including fluorescent proteins expressed in cells.

The design of our DNA-origami structures for generating brightness standards is based on the well-documented 6 helix-bundle (6hb) nanotube, which is ~7 nm in diameter and ~407 nm long^{15,20-22} (Figure 1a & S1–S4). Previously, 6hb and other rod-shaped DNAorigami structures have been used to construct fluorescence markers and barcodes for bioimaging,^{12,22–26} owing to their robust assembly (up to 90% yield) as well as their thermal (melting temperature $\approx 60^{\circ}$ C)²⁷ and mechanical stability (persistence length>1 µm).²⁸ In these applications, single-stranded extensions, termed handles, are placed at designated positions on the surface of DNA nanotubes with nanometer precision to host a myriad of fluorophores. In this work, handles are precisely positioned at 42 bp or ~14 nm apart on each helix to maximize labeling density while minimizing self-quenching. As such, each 6hb structure can accommodate up to 100 copies of fluorophores of interest (e.g., monomeric enhanced green fluorescent protein, mEGFP) in the center, with 12 additional handles at each end reserved for other fluorophores with distinct emission spectra (e.g., Alexa Fluor 647) to aid focusing and quality control. Additionally, 4 handles are evenly spaced on one of the helices to display biotin labels for surface attachment. We designed five versions of 6hb structures to carry 5, 25, 50, 70, and 100

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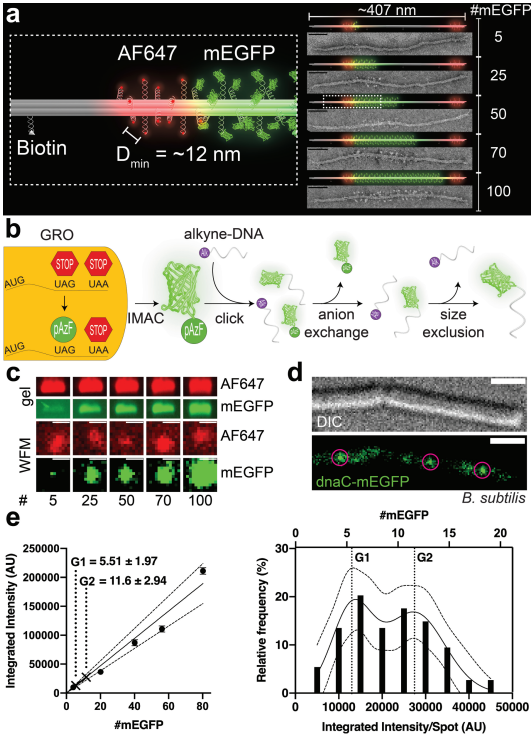


Figure 1. A DNA-origami-based mEGFP brightness standard. (a) 3D models and TEM micrographs of monomeric DNA 6hb structures labeled with 5–100 copies of mEGFP (green) in the main body, 12 copies of Alexa Fluor 647 (red) at each end, and 4 biotin molecules along one side. The minimum spacing of the fluorophores is ~12 nm. Scale bars: 50 nm. (b) Generation of mEGFP-DNA conjugate. mEGFP-pAzF is expressed and purified from a GRO, in which the antisense TAG codon has been reassigned to encode pAzF, an azide-modified Phe. mEGFP(pAzF) was purified *via* immobilized metal affinity chromatography (IMAC), then reacted with alkyne-labeled DNA. Two subsequent purification steps removed unreacted proteins and DNA. (c) Gel electrophoresis (top) and widefield microscopy images (WFM, bottom) of mEGFP standards. Images are set to the same brightness scale (no saturated pixels in the original images). Scale bars: 2 μ m. (d) Differential interference contrast (DIC, top) and WFM (bottom) images of *B. subtilis* (stain NW001) expressing dnaC-mEGFP. Circles indicate puncta picked for quantification. Scale bars: 2 μ m. (e) Quantifying dnaC-mEGFP. Left: a calibration curve with intensities of DNA-origami standards and interpolated protein counts (mean±SEM) from dnaC-mEGFP puncta. Dotted lines denote 95% confidence interval. Right: frequency distribution and sum-of-two-Gaussians fit of dnaC-mEGFP puncta.

mEGFPs (**Figure 1a**), which we prepared from a 7308-nt long circular ssDNA and 5 different pools of synthetic oligonucleotides following well-established DNA-origami assembly and purification protocols.^{16,29–32}

To generate DNA-conjugated mEGFP, we used a Genomically Recoded Organism (GRO) to express mEGFP with a single azide-bearing nonstandard amino acid, pazidophenylalanine (pAzF), at the C-terminus.^{33,34} This was subsequently conjugated to an alkyne-labeled DNA oligonucleotide with complementary sequence to the handles (termed antihandles) by copper-mediated azide/alkyne click chemistry. The conjugation product was purified by anion-exchange and size-exclusion chromatography in two consecutive steps (Figure 1b & S5–S8, see online methods). This conjugation method has advantages compared to conventional crosslinking chemistry such as thiolmaleimide, amine-NHS ester, or SNAP-benzylguanine, in that it cleanly allows 1:1 DNAprotein conjugation with a site-specific, single amino-acid addition that keeps the protein structure and biochemistry perturbation to a minimum. Further, it may be applied to any protein expressed by such GROs using readily available chemicals in aqueous solutions, allowing the creation of imaging standards with any fluorescent protein. The purified mEGFP-antihandle conjugate was then hybridized to the 6hb structures bearing 5-100 handles to generate desired mEGFP standards, which were purified by polyethylene glycol fractionation.³⁵ These mEGFP-labeled structures were first characterized by a quantitative electrophoresis analysis. Each structure's band mobility corresponded well to the designed numbers of mEGFP per structure, with band intensities increasing proportionally to the number of fluorophores (Figure 1c & S9). showing no evidence of mEGFP self-quenching (Figure S10). Negative-stain TEM imaging produced striking micrographs of these decorated structures (Figure 1a & S11) that further confirmed the expected mEGFP density and location on the 6hb tubes.

To demonstrate the utility of our origami-based brightness standards, we used them to quantify dnaC in *B. subtilis*. DnaC is a well-studied helicase that has been shown to assemble into a homo-hexameric ring at the replication fork.^{36–39} Because bacteria contain a single chromosome with two replication forks, dnaC-mEGFP puncta should appear to have 6 or 12 monomeric dnaC, depending on the proximity of the replication forks.³⁹ Cells expressing dnaC-mEGFP and each of the origami standards were immobilized on separate agar pads and imaged with a widefield fluorescence microscope under the same conditions. After subtracting background fluorescence from agar pads and cell autofluorescence, spots were then picked using the ImageJ plugin MicrobeJ⁴⁰ (Figure 1c-e & S12-14). To reduce imaging artifacts, we selected only DNA-origami spots that coincided with slightly elongated Alexa Fluor 647 spots, and dnaC spots that resided within the rod-shaped cells. Spots from origami structures were used to create a standard curve correlating fluorescence intensity to molecule number, which showed excellent linearity (Figure 1e), similar to the bulk measurement by gel electrophoresis (Figure S9). The variance of brightness from the standard structure is consistent with the heterogeneity of fluorescent output of mEGFP molecules⁴¹ and mEGFP labeling efficiency. In order to normalize for sub-stoichiometric labeling of DNAorigami structures, stepwise photobleaching was performed on a 6hb tube designed to carry 5 molecules of Alexa Fluor 488. Fitting the step counts to a binomial distribution

yielded the fluorophore attachment probability of ~0.80 (**Figure S15**); e.g., 100× mEGFP standard had an average of 80 fluorescent proteins, 70× standard had 56, and so on. The labeling efficiency we measured is consistent with previous reports.^{23,42} Finally, the distribution of intensities from bacterial mEGFP puncta were fit to a sum of two Gaussians and calibrated against the standard curve to derive the dnaC stoichiometry. Our method resulted in 5.50±1.97 and 11.6±2.94 (mean±SD) dnaC per puncta (**Figure 1e**), which agrees well with the expected dnaC counts of 6 and 12 molecules.³⁹

In addition to quantifying widefield images of GFP-tagged protein in bacteria, we sought to demonstrate our system's broad applications by counting dye-tagged clathrin light chain (CLC) molecules in mammalian cells using confocal microscopy (**Figure 2**). During receptor-mediated endocytosis, CLC molecules assemble with adaptors and other regulatory proteins into clathrin-coated pits and plaques, distinguished by size, clathrin number, and dynamics.^{43,44} Coated pits are smaller and more circular than plaques, making it relatively straightforward to distinguish between the two in micrographs. Coated pits assemble into clathrin cages containing varying numbers of triskelia, each comprised of 3 clathrin heavy chains and 3 clathrin light chains.⁴⁵ The reported numbers of CLCs in a single vesicle vary widely between different tissues and methods of estimation.^{44,46,47} The recruitment, assembly and disassembly of membrane-coating clathrin structures are highly dynamic (~45–200-s life-time; 60–100 triskelia, or ~180–300 CLCs, for a mature clathrin-coated endocytic pit or vesicle).⁴⁴

To accommodate more fluorophores, we built dimeric 6hb nanotube structures that can in theory host up to ~300 fluorophores (though a maximum of 200 were used in this study). The dimeric nanotube is designed to have 3 barcoding zones reserved for Alexa Fluor 488 or TAMRA (12 fluorophores per zone) to enable the selection of intact dimer structures and, potentially, sorting after simultaneous acquisition²² (Figure 2a). To quantify Halo-tagged CLC labeled with far-red dye silicon rhodamine (SiR)^{48,49} in live HeLa cells, we labeled our fluorescent standards with the same dye. Similar to the mEGFP-DNA conjugation, SiR-azide was conjugated to alkyne-DNA via coppermediated click chemistry^{34,50,51} and then purified using denaturing polyacrylamide gel electrophoresis (PAGE) (Figure S16, also see online methods). Upon hybridizing SiRlabeled anti-handles to the DNA nanotubes, we analyzed the SiR-labeled standards using agarose gel electrophoresis, which showed SiR intensity increasing linearly with the designed dye numbers and the expected barcoding dye combinations (Figure 2b). The biotinylated SiR standards were then purified by rate-zonal centrifugation, individually immobilized on streptavidin-coated glass-bottom dishes, and imaged under confocal microscope (Figure 2a & S17–22), which confirmed high-quality dimeric structures by fluorescent barcode patterns. Next, we applied these standards to counting SiR-labeled CLCs near the surface-adhering membranes of live HeLa cells (Figure 2c-d, S23). Cells and standards were imaged on the same day under identical conditions, and SiR-fluorescent spots were selected from the background-subtracted images using a custom TrackMate⁵² script (Figure 2d, left & S24). SiR puncta in HeLa cells were manually picked to identify those resembling clathrin-coated pits (roundshaped puncta < $\sim 1 \mu m^2$), and CLC counts per cluster were quantified using our DNA-

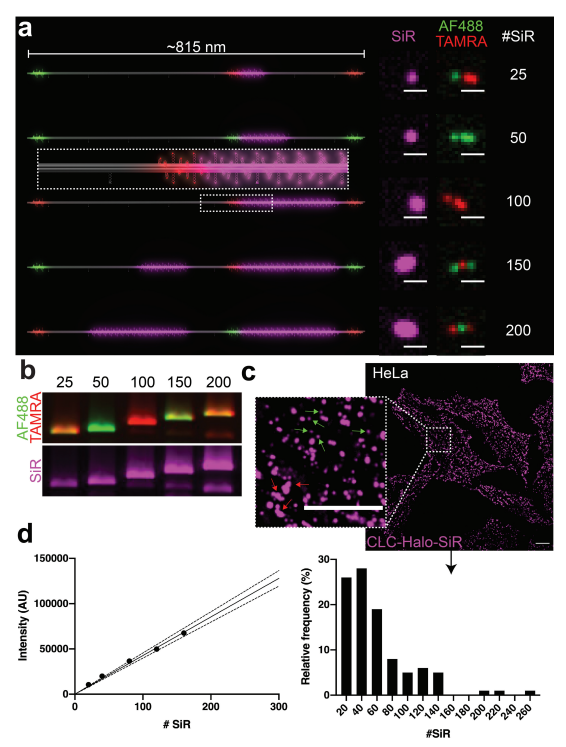


Figure 2. A DNA-origami-based SiR brightness standard. (a) 3D models of dimeric DNA-origami nanotubes hosting 25–200 SiR molecules (magenta) in the main body, as well as Alexa Fluor 488 (Green) and TAMRA (red) at three distinct locations for barcoding. Confocal microscopy images revealed the expected barcoding patterns and corresponding increase in SiR intensity. Scale bars: 1 μ m. (b) Agarose gel images of SiR standards show the expected combinations of barcoding dyes Alexa Fluor 488 and TAMRA, as well as increasing SiR intensity. (c) Confocal images of HeLa cells expressing Halo-fused clathrin light chain (CLC) after labeling with SiR-chloroalkane. Inset shows details of coated pits (small and round, green arrows) and plaques (larger and irregularly shaped, red arrows). Scale bars: 10 μ m. (d) Quantifying SiR-labeled CLCs. Left: Calibration curve generated from DNA-origami-based SiR standards (SEM too small to see). Dotted lines denote 95% confidence interval. Right: Spots containing SiR-labeled CLCs binned by molecule number.

origami calibration curve (**Figure 2d**, right). Out of the 100 clusters, 97 of them contain no more than 150 CLC (median = 47.8), with a decreasing frequency as molecule number increased. This is to be expected from clathrin cages that pinch off shortly after completing assembly.^{44,46} The CLC counts obtained here is likely an underestimate because of the less-than-perfect SiR-labeling efficiency in cells. The few cellular puncta above 150 molecules are presumed to be either accidental selection of clathrin patches, or more than one pit close together.

In summary, our DNA-origami based brightness standards provide a versatile, easy-toimplement system for quantifying 5–300 clustered proteins using fluorescence microscopes readily accessible to cell biology labs. The most valuable features of our standards are their programmability to accommodate fluorophores of various types and stoichiometry, their robustness to withstand near-physiological conditions,⁵³ and their low entry barrier to use without specialized equipment or software. Typically, imaging these standards adds less than 2 hours to a typical cell imaging session (compared to several days using alternative methods), and thus can be easily integrated into biologists' conventional workflows to obtain very good estimates of protein counts. While we demonstrate the standards' efficacy in bacteria and mammalian cells, they should, in principle, be compatible with any cell type. Similarly, because of the large library of DNA-fluorophore conjugation chemistry,⁵⁴ including our GRO-enabled protein conjugation, our system can be used to display and count practically any fluorophore. The primary limitation of our system is that the standards are external to the cell, and so must be imaged in media similar to the cellular compartment being studied. Because the main media effectors of fluorescence output are solvent polarity and pH,⁵⁵⁻⁵⁸ this obstacle can be overcome by matching the pH of the aqueous imaging media to the cellular pH, as is the case in *HeLa* and *B. subtilis* cytosol.^{59,60} In cases where the cellular pH is unknown or dynamic, our universal conjugation method allows attachment of pH-sensitive fluorophores like pHluorin^{55,58} to account for any pH differences. In this work, we used a few existing, commonly used image-analysis software to extract guantitative data from micrographs. However, we envision that a specialized particletracking and pattern-recognition software, perhaps with future development of machine learning, will further expedite the workflow.

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