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Tuning the Sensitivity of Genetically Encoded Fluorescent Potassium Indicators through Structure-Guided and Genome Mining Strategies

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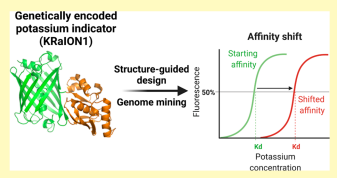
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ABSTRACT: Genetically encoded potassium indicators lack optimal binding affinity for monitoring intracellular dynamics in mammalian cells. Through structure-guided design and genome mining of potassium binding proteins, we developed green fluorescent potassium indicators with a broad range of binding affinities. KRaION1 (K⁺ ratiometric indicator for optical imaging based on mNeonGreen 1), based on the insertion of a potassium binding protein, Kbp, from *E. coli* (Ec-Kbp) into the fluorescent protein mNeonGreen, exhibits an isotonically measured $K_{\rm d}$ of 69 \pm 10 mM (mean \pm standard deviation used throughout). We identified Ec-Kbp's binding site using NMR spectroscopy to detect protein—thallium scalar couplings and refined the structure of Ec-



Kbp in its potassium-bound state. Guided by this structure, we modified KRaION1, yielding KRaION1/D9N and KRaION2, which exhibit isotonically measured K_d 's of 138 \pm 21 and 96 \pm 9 mM. We identified four Ec-Kbp homologues as potassium binding proteins, which yielded indicators with isotonically measured binding affinities in the 39–112 mM range. KRaIONs functioned in HeLa cells, but the K_d values differed from the isotonically measured case. We found that, by tuning the experimental conditions, K_d values could be obtained that were consistent *in vitro* and *in vivo*. We thus recommend characterizing potassium indicator K_d in the physiological context of interest before application.

KEYWORDS: potassium sensor, protein engineering, rational design, NMR structure, genome mining, fluorescent protein

Potassium ions serve key physiological roles in a range of cells and systems, including neurons and glia, ¹⁻⁶ cardiomyocytes, ⁷ and renal cells, ^{8,9} and potassium homeostasis is dysregulated in conditions including seizures, immune cell impairment, and ischemic events. ¹⁰⁻¹² Potassium concentration in cells, tissues, and organisms has been measured using electrodes or dyes. ¹³⁻¹⁵ Recent attention has focused on genetically encoded fluorescent indicators because they could be targeted to specific cells. ¹⁶

The potassium binding protein, Kbp, from *E. coli* (here *E. coli*-Kbp or Ec-Kbp for short) undergoes large conformational changes during potassium binding. Ec-Kbp consists of ~150 amino acids forming two domains, LysM (lysin motif) and BON (bacterial OsmY and nodulation), and binds to one potassium ion.¹⁷ When binding occurs, Ec-Kbp's conformational change brings the two domains closer together and orders the N-terminus such that it lies close to the C-terminus. However, Ec-Kbp's binding site was not previously identified.¹⁷ The conformational change behavior parallels that of calmodulin and M13 in commonly used genetically encoded calcium indicators. Ec-Kbp's conformational change makes it a good candidate for incorporation into genetically encoded

sensors, particularly using split-fluorescent proteins, as used in the calcium indicator NCaMP7.

The protein engineering community has used Ec-Kbp to develop genetically encoded K^+ sensors. Two FRET sensors, GEPII 1.0 and KIRIN1, and a single-fluorescent protein K^+ sensor, GINKO1, have been published. These sensors change fluorescence emission upon K^+ binding, and although they have been used to measure K^+ concentration in solution and cell culture, their binding affinities for K^+ are not appropriate for accurate measurements of intracellular potassium concentrations (previously reported K_d values range from 0.40 to 2.6 mM), whereas textbook measurements of K^+ concentrations are \sim 140–150 mM for mammalian cells at resting potential. Extracellular K^+ concentrations can

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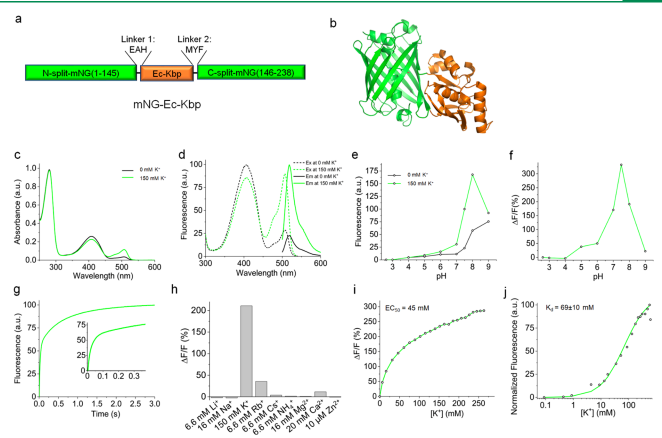


Figure 1. Molecular design and properties of the genetically encoded green potassium indicator KRaION1 in solution. (a) Molecular design of the KRaION1 indicator. (b) The proposed structure of KRaION1 shown as a ribbon diagram according to the crystal structure of NCaMP7 (PDB: 6XW2) and NMR structure of the Ec-Kbp domain (PDB: 5FIM). (c) Absorbance spectra of KRaION1 at 0 and 150 mM potassium at pH = 7.4. (d) Fluorescence spectra of KRaION1 at 0 and 150 mM potassium at pH = 7.4. (e) Relative fluorescence intensity of KRaION1 at 0 and 150 mM potassium as a function of pH. (f) Fluorescence changes upon addition of 150 mM potassium as a function of pH at constant ionic strength. (g) Potassium association using stopped-flow fluorimetry. Association kinetics curves were acquired at a 40 mM final K⁺ concentration starting from K-free protein solution. Inset: the same association kinetics curve shown in the range of the 0–300 ms time frame. (h) Selectivity of KRaION1 to K⁺ and other cations represented as $\Delta F/F$ % measured by titration of Li⁺, Na⁺, K⁺, Rb⁺, Cs⁺, NH₄⁺, Mg²⁺, Ca²⁺, and Zn²⁺. Calculated fluorescence was obtained at the highest expected intracellular concentration for each ion. (i) Fluorescent dynamic range and EC₅₀ of KRaION1 measured in the range of 0–260 mM K⁺. (j) Potassium titration data points (open circles) measured at pH = 7.3 and constant ionic strength of 700 mM, fitted using $Q = (Q_{max} - Q_0)Y + Q_0$ function (green line).

fluctuate up to 30-80 mM in events of sustained neuron depolarization. ²³⁻²⁵ In glia, intracellular K⁺ concentration can increase up to $\sim 60\%$ during potassium concentration regulation. ^{26,27} Without the ability to measure K⁺ dynamically in a cell-specific way, we will not understand K⁺ dynamics in different cell types in healthy and disease states. Our goal was to design and generate intracellularly expressed potassium sensors sensitive to changes in K⁺ comparable to those found in the intracellular milieu.

We report a family of green potassium indicators generated via two approaches. One approach was based on knowledge of the Ec-Kbp K⁺ binding site identified by NMR, while the second approach consisted of identifying and using alternative potassium binding proteins as sensing moieties. Guided mutagenesis was used to tune the indicator mNeonGreen (mNG)-Ec-Kbp's $K_{\rm d}$, yielding two mutants with isotonically measured binding affinities to K⁺ in the range of 96–138 mM and fluorescence dynamic ranges of ~200% over the range of 0.1–150 mM K⁺. The identification of previously unannotated proteins from metagenomic databases that share 45–72% amino acid identity with Ec-Kbp and fusing these to mNeonGreen yielded indicators with isotonically measured

 K^+ binding affinities ranging from 39 to 112 mM and fluorescence dynamic ranges of up to ~200% over the range of 0.1–150 mM K^+ . The indicators generated are ratiometric by excitation. When expressed in HeLa cells that were permeabilized with valinomycin and CCCP ionophores, all tested indicators had K_d values below those measured in isotonic conditions *in vitro*. Thus, K_d could vary depending on the conditions used for its measurement. These indicators thus may be regarded as remaining in the prototype stage, as discrepancies in binding affinities require thoughtful consideration of how K_d is affected by cellular and environmental contexts. Future work is needed to identify the proper calibration conditions in each cell type of interest for the indicators described here and elsewhere.

■ RESULTS AND DISCUSSION

Development of a Genetically Encoded Green Potassium Sensor. We used a design approach based on the insertion of the sensing moiety into a fluorescent protein as has been done with other indicators for calcium: ncpGCaMP6s, NTnC, and NCaMP7. 18,29,30 NCaMP7, which contains mNG as a green fluorescent reporter, is 1.7-fold

Table 1. Spectral and Biochemical Properties of Green Fluorescence Potassium Sensors in Solution and in Cells

	GINKO1		KRaION1		KRaION1/D9N		KRaION2	
	0 mM K ⁺	150 mM K ⁺	0 mM K ⁺	150 mM K ⁺	0 mM K+	150 mM K ⁺	0 mM K ⁺	150 mM K ⁺
absorbance (nm) excitation (nm)	399/501 399/500	400/500 399/500	407/509 408/507	407/509 407/507	409/508 406/506	407/506 407/504	402/508 403/506	404/508 405/506
emission (nm)	514	515	519	518	517	517	516	517
extinction coefficient $(M^{-1} cm^{-1})$	47 000/19 000	46 000/28 000	65 300/12 000	45 600/23 200	42 300/23 700	40 300/20 200	58 400/9300	47 400/26 100
quantum yield (%)	n.d.	7.7/59	35/26	29/57	28/14	23/47	26/23	29/61
pK_a	7.5	7.3	7.7	7.4	7.7	7.3	7.7	7.1
			GINKO1		KRaION1 KRaION1/D9N		KRaION2	
$K_{\rm d}~({\rm mM})^b$		17 ± 7		69 ± 10	138 ± 21		96 ± 9	
$K_{\rm d}~({ m mM})^c$		21 ± 10		42 ± 10	100 ± 6		66 ± 9	
$ au_{ m fast}/ au_{ m slow}~(m ms)^d$		34(85%)/54		31(55%)/71	37(69%)/1211		26(60%)/890	
$\Delta F/F_{ m max}$ (%) e		29		302	434		318	
$\Delta R/R_{\rm max} \ (\%)^f$		n.d.		755 ± 88	1112 ± 160		943 ± 310	
intracellular K_d (mM) ^g		n.d.		37 ± 9	95 ± 21		58 ± 6	

 aK_d was calculated using the same equation used for *in vitro* calculations of K_d . n.d. stands for not determined. $^bK^+$ titration was conducted in isotonic conditions in a range of 0.1-700 mM K⁺. $^cK^+$ titration was conducted at constant 5 mM Na⁺ with an increasing concentration of 0.1-250 mM K⁺. The equation used to fit the titration data and obtain K_d is $Q = (Q_{\text{max}} - Q_0)Y + Q_0$. $^d\tau_{\text{fast}}$ and τ_{slow} indicate time constants with the fluorescence trace fit according to $\Delta F/F(t) = Ae^{-t/\tau_{\text{fast}}} + Be^{-t/\tau_{\text{slow}}} + C$ with the percentage indicating A/(A+B). ^eFluorescence dynamic range $(\Delta F/F_{\text{max}})$ values represent the percent max fluorescence change within the range of 0.1-700 mM K⁺. ^fDynamic range of the excitation fluorescence ratio measured in live HeLa cells calibrated with valinomycin and CCCP within the range of 0-150 mM K⁺. ^gIntracellular characterization of K_d measured in HeLa cells calibrated with valinomycin and CCCP in the range of 0-150 mM K⁺.

brighter than the calcium indicator GCaMP6s¹⁶ and has a maximum $\Delta F/F$ of 2700%. Our potassium indicator design used NCaMP7 as a template, as mNG is brighter than other characterized monomeric GFP proteins³¹ and the design allows for direct insertion of the binding domain into the fluorescent protein.

We replaced NCaMP7's binding moiety with Ec-Kbp while preserving the original amino acid linkers between the fluorescent protein and the binding domain (Figure 1a,b). We performed detailed characterization of mNG-Ec-Kbp's spectroscopic and biochemical properties at 0 and 150 mM under isotonic conditions (Table 1). The absorbance spectrum of mNG-Ec-Kbp shows two peaks at 408 and 508 nm that decrease and increase upon potassium administration, respectively (Figure 1c).³² Correspondingly, mNG-Ec-Kbp's excitation fluorescence spectrum has two peaks at 407 and 507 nm, which result in identical emission maximums at 518 nm (Figure 1d). mNG-Ec-Kbp's extinction coefficient (EC) ratio was 65 300/12 000 M⁻¹cm⁻¹ when excited at 408/507 at 0 mM K⁺ and 45 600/23 200 M⁻¹cm⁻¹ when excited at 407/507 at 150 mM K⁺. The corresponding quantum yields (QYs) obtained at 0 and 150 mM K⁺ when excited at 407/507 nm were 35/26% and 29/57%, respectively (Table 1). Therefore, upon potassium administration, the fluorescence excited at 507 nm increases due to both increasing QY and EC, while fluorescence excited at 407 nm decreases due to both decreasing QY and EC, confirming mNG-Ec-Kbp as a ratiometric indicator, which we named KRaION1: K+ ratiometric indicator for optical imaging based on mNeon-Green 1.

We assessed the pH sensitivity of KRaION1, which exhibits a large fluorescence change in response to 150 mM K⁺ within the pH range of 7–8. The maximum dynamic range ($\Delta F/F$) of 332% was seen at pH 7.5 (Figure 1e,f). Outside this range, the fluorescence response of the indicator decreased. This suggests KRaION1 would be sensitive to fluctuations in pH, as seen for other single fluorescent protein-based indicators. ^{18,29,30} Monitored at pH = 7.4, KRaION1 displayed a fluorescence response

upon binding K+ (at final [K+] = 40 mM) that was biexponential with time constants of τ_{fast} = 31 ms (55% of total amplitude) and $\tau_{\text{slow}} = 71 \text{ ms}$ (remaining amplitude) (Figure 1g). Possible explanations for this include: at the single molecule level, potassium binding to Ec-Kbp's binding pocket is fast, but the previously described 17 conformational rearrangement of the protein is slow; alternatively, there could be two KRaION1 populations: one that can fold rapidly after potassium binding, and another that has to overcome a significant energy barrier to fold. KRaION1's association kinetics response is comparable to that of GINKO1 under identical conditions, characterized by τ_{fast} = 34 ms (85% of total amplitude) and τ_{slow} = 54 ms (remaining amplitude). The fluorescence response of KRaION1 is about 3-fold faster than NCaMP7, perhaps due to the faster association kinetics of potassium ions with Ec-Kbp.

We also assessed KRaION1's ion selectivity by measuring fluorescence changes obtained from titrations with the following ions: Li+, Na+, Rb+, Cs+, NH4+, Mg2+, Ca2+, and Zn²⁺ along with K⁺ (Figure 1h). Only Rb⁺, which has an ionic radius similar to K+, resulted in significant fluorescence increases of KRaION1. At the highest concentrations seen in mammalian cells, $^{33-39}$ KRaION1's $\Delta F/F$ % for K⁺ is the highest at ~200% relative to all other ions, which remain below 50%. The $\Delta F/F$ % response for Li⁺, Ca²⁺, and Zn²⁺ ions increased above 50% only at concentrations that exceed expected intracellular values (Supplementary Figure 1a). Next, we measured KRaION1's fluorescence response in conditions mimicking intracellular ion compositions⁴⁰ by increasing potassium concentration until fluorescence reached an apparent plateau. The fluorescence increased to ~286% of $\Delta F/F$ at ~250 mM K⁺, and the corresponding EC₅₀ was ~45 mM (Figure 1i).

KRaION1's $K_{\rm d}$ was obtained by fitting the fluorescence response to a single site binding model and was 69 \pm 10 mM under isotonic conditions of variable K⁺ (Figure 1j). For comparison, GINKO1 had a $K_{\rm d}$ value of 17 \pm 7 mM under similar conditions. These $K_{\rm d}$ values are less useful for the

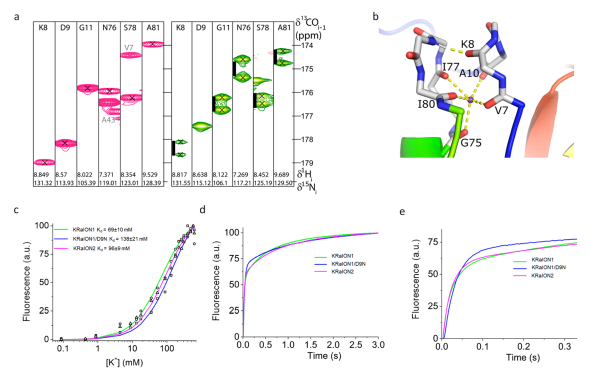


Figure 2. Identification and modification of Ec-Kbp's potassium binding site. (a) 1 H and 13 C strips from HNCO spectra of K⁺ (magenta) and Tl⁺ (green) bound Ec-Kbp showing the residues whose crosspeaks are split by an additional J coupling in the Tl⁺ bound form. The black scale bars represent 100 Hz. (b) Refined configuration of the Ec-Kbp ion binding site. The ion is coordinated entirely by backbone carbonyls from loops 1 and 5 (backbone atoms shown as sticks). Sections of the rainbow-colored cartoon of the rest of the protein are visible. Coordinating carbonyl to K⁺ interactions and the K8 to G79 hydrogen bond are shown as yellow dashed lines. (c) Potassium titration data points for KRaION1 and mutants KRaION1/D9N and KRaION2 (open circles, squares, and triangles, respectively) measured at pH = 7.4 and constant ionic strength, fitted using the equation $Q = (Q_{max} - Q_0)Y + Q_0$ (denoted by a green line, blue line, and magenta line, respectively). (d) Potassium association using stopped-flow fluorimetry. Association kinetics curves were acquired at 40 mM final K⁺ concentration starting from K-free protein solution. (e) Association kinetics curves in the range of the 0–300 ms time frame.

detection of small changes in mammalian cells' intracellular potassium concentration, so we sought to optimize KRaION1 to achieve a higher $K_{\rm d}$. To achieve this, we sought additional information about Ec-Kbp's ion binding site.

Identification of Ec-Kbp's Ion Binding Site. We used NMR spectroscopy to locate Ec-Kbp's ion binding site. In NMR spectroscopy, local chemical shift differences between free and ligand-bound forms of proteins often provide evidence of binding site locations, but this was not feasible for Ec-Kbp due to the extreme line-broadening of many signals in its free form. Evidence for potassium binding site locations can be obtained using ¹⁵N-labeled ammonium ions to substitute for K^{+,41,42} However, Ec-Kbp unfolds at the pHs at which the ammonium ¹H signal is distinct from the water ¹H signal. Isotopes of potassium are quadrupolar nuclei with modest sensitivity, making them unsuitable for determining proteinion contacts in solution. Instead, we used thallium(I) as a K+ mimic, allowing the measurement of direct two bond scalar couplings (${}^{2}J^{-13}C=0...^{205/203}Tl^{+}$) (B. O. Smith, in preparation) of around 100 Hz in the HNCO spectrum (Figure 2a) that unambiguously identify the backbone carbonyls of V7 and A10 from loop 1 in Ec-Kbp's N-terminal extension and G75, 177, and 180 from loop 5, the turn immediately preceding the BON domain's last β -strand, as Tl⁺ ligands. We believe this is the first reported observation of scalar (J) coupling between a protein's ¹³C enriched carbonyl carbons and a bound thallium ion. The large J couplings observed provide direct and unambiguous evidence of which groups coordinate with the

ion. In fact, the variation in J for the different residues suggests the data may contain additional information about the interatomic distances and angles that could be decoded in the future.

Refined Structure of Ec-Kbp·K⁺. To provide an accurate structural model on which to base the mutagenesis of KRaION1's potassium binding site, we recalculated the Ec- $Kbp \cdot K^+$ structure 17 by incorporating a K^+ ion restrained to the carbonyl oxygens identified as Tl+ ligands (PDB: 7PVC; Supplementary Table 1). The coordinating carbonyl oxygens were restrained to be between 2.75 \pm 0.1 Å of the K⁺ ion, as suggested by Harding⁴³ without any constraint on the C=O··· K⁺ bond angle. Since the NMR structure calculation software, ARIA2.3/CNS, uses the OPLX force field with explicit TIP3Plike water representation for the water refinement stage, 44 we used the parameters for K+ suggested by Mamatkulov and Schwierz. 45 The refined structure differs little from the structure calculated without an ion included (Supplementary Figure 2a) but binds K⁺ between loops 1 and 5 at the center of a distorted square pyramid whose base of $\sim 3.3-4.7$ Å on each side is formed by the carbonyl oxygens of residues V7, A10, 177, and 180, while G75's carbonyl oxygen sits at the apex with the lateral sides ranging from 3.56 ± 0.40 Å for G75 and I80 to 5.05 ± 0.29 Å for G75 and V7. The five backbone carbonyls coordinate K+ at distances between 2.62 and 2.86 Å (Supplementary Table 2). K8's carbonyl oxygen could be a sixth K⁺ ligand forming the vertex opposite G75's carbonyl oxygen in an elongated octahedral coordination sphere.

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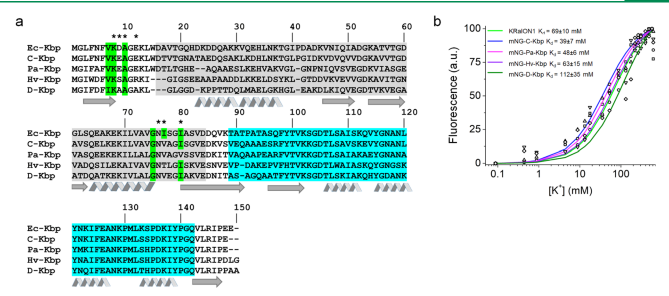


Figure 3. Identification of alternative potassium binding proteins and comparison to Ec-Kbp. (a) Alignment of amino acid sequences of Ec-Kbp with five homologues obtained from a metagenomic BLAST search. Residues comprising the LysM and BON domains are shaded in gray and cyan, respectively. Residues highlighted in green indicate those that are conserved with the Ec-Kbp-identified potassium binding site. The β-sheet-forming regions and α-helix-forming regions are denoted by arrows and ribbons, respectively. Residues that were selected for site-directed mutagenesis in Ec-Kbp are denoted by asterisks. (b) Potassium titration data points for KRaION1 and homologues mNG-C-Kbp, mNG-Pa-Kbp, mNG-Hv-Kbp, and mNG-D-Kbp (open circles, squares, upward triangles, downward triangles, and diamonds, respectively) measured at pH = 7.4 and constant ionic strength, fitted using the equation Q = $(Q_{max} - Q_0)Y + Q_0$ (denoted by a green line, blue line, magenta line, purple line, and dark green line, respectively).

However, it lies more distant from K⁺ (3.98 \pm 0.37 Å) and is instead involved in a hydrogen bond with the backbone amide of G79 (Figure 2b). The conserved glycine residues in the two loops allow the ion binding site to form. G11 adopts an extended conformation with a positive φ dihedral angle, and its lack of side chain allows the close approach of the two loops. G75 is in an α -helical conformation, but the side chain of any other residue at this position would disrupt the close packing of the conserved F6 with its neighboring residues. G79 contributes to a tight turn with a positive φ , allowing its amide to hydrogen bond with K8. The other conserved residue in loop 5, N76, has a role in the overall fold of the protein as its side-chain amide makes a pair of interdomain hydrogen bonds with V143 in the LysM domain. As well as contributing to ion binding, A10 likely ensures close contact can be made with V7 and G141 at the N-terminus of the final beta-strand in the LysM domain.

Pentavalent coordination of potassium in proteins is commonly observed, ⁴⁶ but it is possible that a water molecule could be involved in coordinating the ion raising the coordination number. The structure reveals a likely explanation for Ec-Kbp's specificity for potassium and slightly larger ions over sodium: the two ion binding loops are already tightly packed with potassium bound, while accommodating sodium would dictate a shortening of each of the ion—carbonyl distances by approximately 0.36 Å to maintain favorable interactions, which does not seem sterically feasible. The structure also explains the roles of most of the conserved residues surrounding the ion binding site and explains why it is difficult to find mutations that subtly alter potassium affinity without abolishing it altogether.

Structure-Guided Mutagenesis Modulates K⁺ **Binding.** To fine-tune Ec-Kbp's potassium binding affinity, we introduced mutations within and around the ion binding site, informed by the new NMR structure (Figure 2b and

Supplementary Figure 2b). Since K⁺ appears to be coordinated by the protein's backbone carbonyl groups, our scope for directly affecting the binding site was limited. We speculated that adjusting the electrostatic character of residues near the binding pocket could affect the stability of the K⁺ bound state and increase the K_d. We generated a set of KRaION1 mutants with altered residues within and near the K+ binding site. K8, A10, and N76 were swapped for another small residue (A, G) or to a negatively charged D. These choices were made on the basis of the residues' electrostatic characteristics: positive or neutral residues were swapped for other neutral or negative residues, respectively. The conserved hydrophobic residues I77 and I80 were mutated to smaller amino acids that are either neutral, like A and G, or small and polar, like S. We deduced that nearby residues D9 and E12 could be mutated to remove their negative charges and make the local electrostatic environment less favorable for K⁺ binding without disrupting binding altogether. These residues were substituted with either polar, positively charged, or small nonpolar residue groups: N/ A/K or Q/A/K.

We measured the fluorescence and absorbance spectra of the KRaION1 mutants over a range of K⁺ concentrations from 0 to 310 mM under nonisotonic conditions at pH = 7.3. We determined their apparent $K_{\rm d}$, $\Delta F/F$, baseline fluorescence brightness, and protein folding efficiency (Supplementary Table 3). We used nonisotonic conditions for quick screening of the variants' functionality but used isotonic conditions for proper assessment of $K_{\rm d}$ to remove the effects of varying ionic strength on our measurements. KRaION1's apparent $K_{\rm d}$ in these nonisotonic conditions was 12 \pm 4 mM in comparison to a $K_{\rm d}$ of 69 \pm 10 mM in isotonic conditions, suggesting the indicator's $K_{\rm d}$ is affected by ionic strength. Variants with mutations made directly to the K⁺ binding site appeared to have disrupted functioning, as we observed fluorescence dynamic ranges of <30% and folding efficiency of <50%

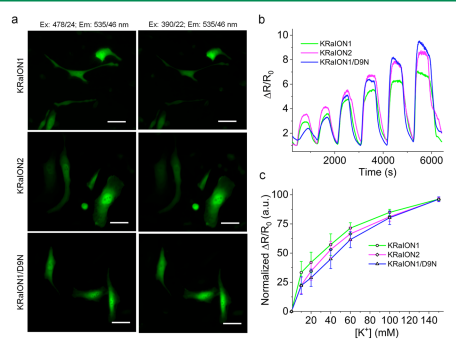


Figure 4. Calibration of KRaION1, KRaION2, and KRaION1/D9N in live HeLa cells. (a) Representative fluorescence images of KRaION1, KRaION2, and KRaION1/D9N in the green channel with UV and cyan excitations (n = 10 FOVs from three independent transfections each). Scale bars, 10 μm. (b) Representative excitation ratiometric response of KRaION1, KRaION2, and KRaION1/D9N at increasing [K⁺] from 0 to 150 mM, equilibrated with valinomycin and CCCP (n = 16, 18, and 21 cells, respectively, from 3 independent transfections; ratio was calculated as $F_{478 \text{ nm}}/F_{390 \text{ nm}}$ for the spectral configuration shown in (a)). (c) Binding titration curves for KRaION1, KRaION2, and KRaION1/D9N when equilibrated with valinomycin and CCCP (n = 15, 20, and 20 cells from 2 independent transfections; KRaION1, and KRaION2, and KRaION1/D9N, respectively; data points, mean; error bars, standard deviation) presented as normalized excitation ratio $F_{478 \text{ nm}}/F_{390 \text{ nm}}$. Experiments were done at 33 °C.

relative to KRaION1. However, mutants D9N and E12A, residues adjacent to the K+ binding site, exhibited higher isotonically measured $K_{\rm d}$ values of 138 \pm 21 and 96 \pm 9 mM with a $\Delta F/F_{\text{max}}$ of 434% and 318%, respectively (Figure 2c). This suggested that changing the electrostatic environment of the binding pocket is possible through structure-guided rational design. Both mutants maintained selectivity to K⁺ over other ions and showed a decreased $\Delta F/F$ % response to Zn²⁺ (Supplementary Figure 1b,c). Relative to KRaION1, both mutants had different biexponential kinetics of association to K⁺. D9N had activation time constants of τ_{fast} = 37 ms (69% of total amplitude) and τ_{slow} = 1211 ms (remaining amplitude) (Figure 2d), and E12A had activation time constants of τ_{fast} = 26 ms (60% of total amplitude) and $\tau_{\rm slow}$ = 890 ms (remaining amplitude) (Figure 2e). The D9N mutation led to slower activation kinetics, whereas the E12A mutation led to faster activation kinetics relative to KRaION1 and GINKO1. We named the mNG-Ec-Kbp-E12A mutant KRaION2 because of its favorable K_d and kinetics values.

Ec-Kbp Homologues Can Be Used as Alternative Binding Moieties. As an alternative approach for indicator design with a binding affinity relevant to the intracellular mammalian environment, we searched for Ec-Kbp homologues to use as the sensing moiety in our sensor. We used protein BLAST to search through a data set (env_nr) that contains proteins from whole genome shotgun sequencing (WGS) metagenomic projects. We selected four proteins with unknown functions that shared 45–72% amino acid identity with Ec-Kbp and in which the K⁺ binding site was also largely conserved (Supplementary Table 4, Figure 3a). Protein homologue names are based on the organism or origin of sample collection: Ec, *E. coli* (NCBI reference sequence:

WP_000522415.1); C, compost (GenBank: MNG82101.1); Pa, P. aeruginosa (NCBI reference sequence: NP_253865.1); Hv, hydrothermal vent (GenBank: VAV91021.1); D, Defluviicoccus sp. (GenBank: SUS08588.1). Amino acid changes observed in the putative potassium binding site maintained the original residues' electrochemical properties. A change in binding site residue V7 to I7 was observed in D-Kbp. Changes to residue I77 to V at equivalent positions were seen in all identified homologues with the exception of Hv-Kbp whose residue change was to T. Another change was seen in the Pa-Kbp homologue, in which residue I80 was swapped for V.

We exchanged Ec-Kbp for each of these homologues in the KRaION1 indicator to check their functionality. When tested at different K^+ concentrations under isotonic conditions, all sensors had a fluorescence response that varied with K^+ concentration, suggesting these homologues are also K^+ binding proteins and sequence changes near the binding site did not disable their K^+ binding function (Supplementary Table 5, Supplementary Figure 3). These alternative binding moieties confer a range of isotonically characterized binding affinities to the indicators from 39 to 112 mM (Figure 3b). One homologue, mNG-D-Kbp, displayed an isotonically measured $K_{\rm d}$ value of 112 \pm 35 mM in the same range as KRaION1 and its D9N and E12A variants. Thus, alternative potassium binding domains can generate promising potassium indicator candidates with a variety of $K_{\rm d}$'s.

In Vivo Characterization of Indicators in HeLa Cells. We focused on a subset of indicators to characterize in cultured HeLa cells: KRaION1, KRaION2, and KRaION1/D9N. In addition to KRaION1, we chose the D9N and E12A mutants due to their higher K_d , fast kinetics, and lower sensitivity to Zn^{2+} . We excluded mNG-D-Kbp from this

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experiment because of its lower $\Delta F/F$ %. We used UV and cyan light to excite the fluorescence of KRaION1, KRaION2, and KRaION1/D9N (Figure 4a). To permeabilize the cells and allow intracellular diffusion of potassium while maintaining steady pH, we used the ionophores valinomycin and CCCP. We exchanged different concentrations of K⁺ into the extracellular space, and in between exchanges, cells were incubated in 0 mM K⁺ to allow sensor equilibration to baseline fluorescence. The ratio of fluorescence emission changes $(F_{487 \text{ nm}}/F_{390 \text{ nm}})$ showed KRaION1, KRaION2, and KRaION1/D9N responses when pulsed with K+ buffers ranging from 0 to 150 mM K+, suggesting the indicators are still functional in cells (Figure 4b). The excitation fluorescence ratio of the indicators measured when excited at two distinct wavelengths, $\Delta R/R_0$, increased up to 150 mM intracellular K⁺ with the max $\Delta R/R_0$ at 755 \pm 88%, 943 \pm 310%, and 1112 \pm 160% for KRaION1, KRaION2, and KRaION1/D9N, respectively (Table 1).

We calculated the sensors' K_d values when expressed in HeLa cells as 37 \pm 9 mM for KRaION1, 58 \pm 6 mM for KRaION2, and 95 \pm 21 mM for KRaION1/D9N (Figure 4c). We selected KRaION1/D9N as the indicator with the highest fluorescence ratio to evaluate whether cells with a high expression of indicator have a different K_d value than those with lower expression. K_d values obtained accounting for these expression differences were 90 \pm 41 and 94 \pm 14 mM for high and low expressing cells, respectively, which overlap with our previously reported measurement that assumed equal expression across cells (Supplementary Figure 4a). However, these K_d values are inconsistent with the isotonically measured K_d values obtained in vitro. Cell calibration experiments performed with valinomycin alone without CCCP were even more inconsistent (Supplementary Table 6, Supplementary Figure 4b). When performing the same calibration experiment with valinomycin for GINKO1, we obtained an intracellular K_d value of 3 \pm 0.23 mM, different from its isotonic value.

We attempted to understand this K_d discrepancy by measuring the indicator's binding affinity at constant 5 mM Na⁺ and variable K⁺ in the range of 0.1 to 250 mM. In these conditions, we obtained a different K_d value for KRaION1 of 42 ± 10 mM, lower than the isotonic case but overlapping the HeLa cell case when calibrated with valinomycin and CCCP (Table 1). Overlap in K_d values between this in vitro measurement and cell calibration was also observed for KRaION2 and KRaION1/D9N at 66 ± 9 and 100 ± 6 mM, respectively. Different K_d values were observed in all other measured indicators under these in vitro conditions (Supplementary Table 7). This suggests the indicators are sensitive to changes in ionic strength when measured in vitro. K_d discrepancies have previously been studied for calcium and potassium probes. ^{28,47} For potassium probes, a combination of several ionophores and sucrose was most effective in potassium calibration experiments in Jurkat and U937 cell types. 28 This is consistent with our observations of cell calibration with a combination of valinomycin and CCCP versus just valinomycin. For these calibration experiments, we suggest identifying the best combination of ionophores for proper equilibration of ion concentrations in cell types of interest. Future potassium indicator engineering efforts may benefit from screens being fully performed in cell types of interest rather than in vitro with methods like robotic molecular screening, which has not yet been applied to potassium indicators.⁴⁸

CONCLUSION

We have designed a set of potassium indicators on the basis of the insertion of Ec-Kbp mutants and homologue variants into mNeonGreen. Insertion of Ec-Kbp into mNeonGreen results in KRaION1, a bright and fast indicator, with an isotonically measured binding affinity for potassium of 69 \pm 10 mM. We used two approaches to increase this K_d value: structure-guided mutagenesis and identification of alternative potassium binding proteins. We utilized NMR to determine the structure of Ec-Kbp to reveal that the backbone carbonyls of residues V7, A10, G75, I77, and I80 coordinate the potassium ion. With this information, we designed two mutants, KRaION1/D9N and KRaION2, with higher isotonically measured K_d values of 138 \pm 21 and 96 \pm 9 mM, respectively. We also identified four Ec-Kbp homologues that have a conserved binding site for potassium and can be alternative binding moieties in the indicator. When expressed intracellularly, KRaION1, KRaION2, and KRaION1/D9N maintain their functionality. However, K_d values measured in the intracellular environment with valinomycin and CCCP were lower than those observed in vitro. The measurement of K_d at constant sodium concentration gave values that overlapped with these intracellular conditions, raising the point that physiological characterization of potassium indicators needs to be performed carefully and may need to occur in the exact cell type and physiological context of interest. KRaION1's apparent K_d for potassium was already more than 2 orders of magnitude higher in vitro than Ec-Kbp's. Perhaps Ec-Kbp's conformational change is frustrated by having its N- and C-termini tethered by mNG. This suggests further optimization of the linker sequences to mNG may be useful in optimizing KRaION1's $K_{\rm d}$. For all indicators, further structure-guided mutagenesis can be combined with other techniques such as high-throughput, multiparameter directed evolution approaches 48 to further improve other parameters of interest, such as brightness and kinetics.

■ MATERIALS AND METHODS

Molecular Cloning and Mutagenesis. De novo gene synthesis of the designed potassium sensor including Ec-Kbp and identified homologues and subsequent subcloning into the pBAD-HisD vector were done by Genscript. The genes were codon-optimized for human cells. Mutagenesis of individual residues in KRaION1 was done using the QuikChange Site-Directed Mutagenesis Kit (Agilent). Forward and reverse primers used to generate each mutant are provided in Supplementary Table 8. Indicators were transformed into TOP10 (Invitrogen) electrocompetent E. coli cells and grown on LB plates with 1:1000 dilution of carbenicillin and 0.002% arabinose. Confirmation of the correct mutation for each construct was done by colony sequencing (Eton Bioscience Inc.). For mammalian expression, the selected indicator genes were cloned into a mammalian expression vector pN1 (Clontech). The DNA sequence of GINKO1 was obtained from Addgene (plasmid #113112).

Protein Purification and *in Vitro* **Characterization.** Proteins were expressed and purified using the pBAD/HisB arabinose-inducible system (Invitrogen) in TOP10 cells (Invitrogen) cultured in RM medium as previously described. ^{49,50} Before measurements, the proteins were dialyzed against various buffers depending on the subsequent measurement conditions (see sections below) and stored at 4 °C.

The following buffers were prepared for characterization: 150 mM NaCl, 25 mM HEPES, 25 mM MES, and 10 mM Tris (to be used as 0 mM $\rm K^+$) and 150 mM KCl, 25 mM HEPES, 25 mM MES, and 10 mM Tris. Absorbance was measured using a spectrophotometer (Hitachi Shimadzu UV-3600 Plus). The extinction coefficients were

determined using alkine denaturation with 1 M NaOH as described previously. Fluorescence was measured using two excitation wavelengths at 400 and 480 nm and emission ranges of 410–700 and 490–700 nm, respectively, in the same buffers with 0 and 150 mM $\rm K^{+}$. Quantum yields were measured at 0 and 150 mM $\rm K^{+}$ buffers at excitation wavelengths of 405 and 480 nm (Edinburgh FLS1000 spectrometer).

For p K_a measurements, 0 and 150 mM K⁺ buffers at various pH values were used to measure the fluorescence intensity in the range of 250–600 nm (Plate Reader Thermo Varioskan LUX). The kinetics of the fluorescence response was obtained with 480 nm excitation and 520/40 nm emission at 40 mM K⁺ concentration using stopped flow spectrophotometry (Applied Photophysics Ltd., SX 20).

For K⁺ titration under nonisotonic conditions, purified proteins in artificial mammalian cell cytoplasm buffer (12 mM NaHCO₃ and 1 mM MgCl₂ in 25 mM Tris/MES, pH 7.4) were diluted into a series of buffers with K⁺ (potassium D-gluconate) concentrations ranging from 0 to 260 mM. For the examination of K⁺ specificity, purified proteins were diluted into a series of buffers that each contained a different salt, these being: LiCl, NaCl, KCl, RbCl, CsCl, NH₄Cl, MgCl₂, and CaCl₂ ranging from 0 to 260 mM and ZnCl₂ at a range of 0 to 260 μM. The fluorescence spectrum of the purified proteins in each solution was measured with excitation at 490 nm and emission from 510 to 600 nm (Thermo Scientific Varioskan LUX).

For $K_{\rm d}$ characterization under isotonic conditions, the proteins were dialyzed against 25 mM MES, 25 mM HEPES buffer at pH 7.3 and added to a series of isotonic solutions with varied KCl to NaCl ratios. KCl concentrations tested were in the range of 0 to 700 mM KCl. For characterization of $K_{\rm d}$ in a secondary condition, the 25 mM MES, 25 mM HEPES buffer at pH 7.3 with constant 5 mM NaCl concentration and increasing concentrations of KCl in the range of 0.1 to 250 mM K⁺ was used. Absorbance and fluorescence from three technical replicates for each indicator were measured at 24–26 °C (Tecan Spark Plate Reader). Titration curves and $K_{\rm d}$ were later obtained by fitting to the following equation using a custom Python script: $Q = (Q_{\rm max} - Q_0)Y + Q_0$ where $Q_{\rm i}$ is fluorescence and $Q_{\rm max}$ and Q_0 are maximum and minimum fluorescent yields, respectively.

$$Y = \frac{([M_{\rm tot}] + [L_{\rm tot}] + K_{\rm d}) \pm \sqrt{([M_{\rm tot}] + [L_{\rm tot}] + K_{\rm d})^2 - 4[M_{\rm tot}][L_{\rm tot}]}}{2[M_{\rm tot}]}, \text{ where } M \text{ is a}$$

fixed concentration component of the genetically encoded sensor, L is the varying concentration component of K^+ , and K_d is the dissociation constant. All calculated values are expressed as K_d \pm standard deviation.

Ec-Kbp·Tl⁺ NMR and Ec-Kbp·K⁺ Structure Determination. To identify the amino acid groups that ligate the cation, a 0.25 mM sample of U– 13 C- and U– 15 N-labeled Ec-Kbp prepared as described previously 17 was diluted in 20 mM sodium phosphate, pH 7.2, 0.01% w/v sodium azide, and 5% D₂O with the addition of 1 mM thallium sulfate (to achieve 2 mM Tl⁺). From a 3D HNCO experiment recorded (1024 × 27 × 128 complex points for sweep widths of 9615 × 2127.7 × 2413.8 Hz in 1 H, 15 N, and 13 C, respectively) at 14.1 T and 298 K on a Bruker AVANCE IIIHD spectrometer equipped with a TCI cryoprobe and processed conventionally, the cross peaks that were split by an additional coupling in the 13 CO dimension were identified, making use of the backbone resonance assignment of Ec-Kbp·Tl⁺ (B. O. Smith, personal communications) in CCPN analysis software. 52

The refined structure of the Ec-Kbp·K⁺ was calculated using the same restraints as in the original structure determination 17 but with the addition of a potassium ion and 5 distance restraints of 2.75 ± 0.1 Å between it and the five coordinating carbonyls. 43 The parameters for the singly charged potassium ion were added to the topallhdg5.3.pro and parallhdg5.3.pro files in the ARIA2.3/CNS software with the epsilon and sigma values set to 0.62 kJ mol $^{-1}$ and 3.0695 Å in the OPLX section used during refinement in explicit water. 45 Eight rounds of structure calculations starting from randomized coordinates in Cartesian space were executed with 100 structures each. The 20 structures from the final iteration with the lowest restraint energy were refined using the ARIA2.3/CNS software in explicit water. The refined structures were analyzed and figures

prepared using PyMOL and Inkscape. The Ec-Kbp· K^+ structure has been deposited at the protein data bank (PDB: 7PVC).

Metagenomic Search for Alternative Ec-Kbp Domains Using Protein BLASTp. To search for alternative potassium binding domains, we used NCBI's BLASTp protein to protein sequence alignment tool. The Ec-Kbp amino acid sequence (NCBI reference sequence: WP_000522415.1) was used to search against the metagenomic database (env_nr). No other parameters were set to perform the search. The obtained results were filtered by percentage similarity of the amino acid sequences and were picked manually. The *E*-value metric was also taken into consideration, where numbers closest to 0 would be identified as good matches.

In Vitro Screening of KRalON1 Mutants and Homologues. Protein expression and purification was done as described above. To screen mutants and homologues for binding affinity, the proteins were dialyzed against 100 mM Tris buffer at pH 7.3 and assessed at different KCl concentration increments ranging from 0 to 310 mM KCl in 100 mM Tris at pH 7.3 using a Tecan Plate Reader (Tecan Spark). Sensor titration curves were fitted, and $K_{\rm d}$ values were obtained fitting into the single site binding equation described above.

In Vivo Characterization and Imaging in HeLa Cells. HeLa cells (ATCC) were cultured in Dulbecco's modified Eagle's medium (Gibco) with 10% fetal bovine serum (FBS; YEASEN Biotech) and were incubated at 37 °C with 5% CO₂. Cells were plated on 12 mm coverslips (Fisher Scientific) coated with Matrigel in 24-well plates before transfection. Liposomal transfection was applied according to the manufacturer's protocol (YEASEN Biotech). HeLa cells were transfected using Hieff Trans (YEASEN Biotech) according to the manufacturer's protocol and imaged 36–48 h post-transfection using an inverted wide-field Nikon Eclipse Ti2 microscope equipped with a SPECTRA III light engine (Lumencor) and a Orca Flash4.0v3 camera (Hamamatsu), controlled by NIS-Elements AR software, and using a 20×, 0.75 NA objective lens.

The solutions with varying concentrations of K⁺ were prepared by mixing two stock solutions (25 mM MES, 25 mM HEPES, and 250 mM NaCl as the "no potassium solution" and 25 mM MES, 25 mM HEPES, and 250 mM KCl with potassium) in the corresponding ratios. Cell medium was replaced with 500 μ L of 25 mM, 25 mM HEPES, and 250 mM NaCl buffer with 15 μ M valinomycin and 5 μ M CCCP and incubated for 15 min before perfusing. The solutions ranging from 0 to 150 mM K⁺ were administered with 15 μ M valinomycin (Aladdin Biochemical Technology Co., Ltd.) and 5 μ M CCCP (Shanghai Macklin Biochemical Co., Ltd.) right before perfusion. Cell calibration was also performed in the same manner with just 15 μ M valinomycin.

To perform automated buffer exchange, the coverslip with the cells was transferred to the RC-26G flow chamber (Warner Instruments, USA) and connected to a custom-built perfusion system consisting of a peristaltic pump (Baoding Chuangrui Precision Pump Co., Ltd.), the SV06 12 port switch value (Runze Fluid, China), controlled heated platform (Warner Instruments, USA), and vacuum pump. The system was designed to offer programmable buffer exchange and continuous monitoring of flow rate, temperature, and channel switching. The perfusion system was controlled by the custom LabVIEW code (National Instruments Corporation; Supplementary Figure 5).

For live cell imaging, the programmable perfusion system was used to provide the series of extracellular buffers at a 1 mL/min flow rate with different potassium concentrations (0 to 150 mM $\rm K^+$ as described) containing valinomycin and CCCP or valinomycin alone, ensuring that the cells were bathed in a consistent environment without pipetting the desired solution manually. The buffer temperature was kept at 33 $^{\circ}\rm C$ throughout the entire imaging.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssensors.1c02201.

NMR and refinement statistics; geometry of the K^+ binding site; physical properties of the tested KRaION1 mutants and homologs; search results obtained from protein BLAST of the Ec-Kbp sequence; binding affinity and dynamic range characterization; properties of indicators measured in HeLa cells; binding affinity (K_d) measurements; primer sequences; selectivity of KRaION1, KRaION1/D9N, and KRaION2 to K^+ ; structure of Ec-Kbp· K^+ and mutation sites; cell calibration curves; experimental workflow of the perfusion system setup (PDF)

Accession Codes

Sequences of the reported proteins are available at Genbank at the following accession codes: KRaION1, OK187805; mNG-Ec-Kbp-D9N (KRaION1/D9N), OK187806; KRaION2, OK187807; mNG-C-Kbp, OK187808; mNG-Pa-Kbp, OK187809; mNG-Hv-Kbp, OK187810; mNG-D-Kbp, OK187811. The solution structure of potassium bound Ec-Kbp has been deposited in the Protein Data Bank, accession code: 7PVC. Plasmids of interest are available from Addgene.

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C.C.T.C., K.D.P., and E.S.B. initiated the project. C.C.T.C., F.V.S., and K.D.P. designed the mNG-Ec-Kbp (KRaION1) indicator. C.C.T.C. and K.D.P. developed the indicators, performed mutagenesis and screening, and identified Ec-Kbp homologues. C.C.T.C. and M.Y. performed *in vitro* characterization and data analysis. C.L. and K.D.P. performed sensor characterization in HeLa cells. L.Y. and K.D.P. performed ion selectivity experiments. B.O.S. determined the Ec-Kbp NMR structure and provided guidance on mutagenesis and data analysis. C.C.T.C., B.O.S., K.D.P., and E.S.B. interpreted the data and wrote the manuscript with contributions of all authors. K.D.P. and E.S.B. oversaw all aspects of the project.

Notes

The authors declare no competing financial interest. We acknowledge Henrietta Lacks and her family for their immense contributions to scientific research. The mammalian cell experiments in this study were conducted on HeLa cells. The HeLa cell line was established from the tumor cells of Henrietta Lacks without her knowledge or consent. Since 1951, Henrietta Lacks has made countless contributions to medicine and science. E.S.B. makes a donation, on behalf of the lab, to the Henrietta Lacks Foundation for each publication the Boyden lab makes that uses HeLa cells, as inspired by the Reck-Peterson lab's example.

The TOC image was created with BioRender.com.

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