An Open Library of Human Kinase Domain Constructs for Automated Bacterial Expression

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Supporting Information

ABSTRACT: Kinases play a critical role in cellular signaling and are dysregulated in a number of diseases, such as cancer, diabetes, and neurodegeneration. Therapeutics targeting kinases currently account for roughly 50% of cancer drug discovery efforts. The ability to explore human kinase biochemistry and biophysics in the laboratory is essential to designing selective inhibitors and studying drug resistance. Bacterial expression systems are superior to insect or mammalian cells in terms of simplicity and cost effectiveness but have historically struggled with human kinase expression. Following the discovery that phosphatase coexpression produced high yields of Src and Abl kinase domains in bacteria, we have generated a library of 52 His-tagged human kinase domain constructs that express above 2 μg/mL of culture in an automated expression screen of 96 human kinase constructs. Using a fluorescence-based thermostability assay and a fluorescent ATP-competitive inhibitor, we show that the highest-expressing kinases are folded and have well-formed ATP binding sites. We also demonstrate that these constructs can enable characterization of clinical mutations by expressing a panel of 48 Src and 46 Abl mutations. The wild-type kinase construct library is available publicly via Addgene.

Kinases play a critical role in cellular signaling pathways, controlling a number of key biological processes that include growth and proliferation. There are more than 500 kinases in the human genome, many of which are of therapeutic interest. Perturbations due to mutation, translocation, or upregulation can cause one or more kinases to become dysregulated, often with disastrous consequences. Kinase dysregulation has been linked to a number of diseases, such as cancer, diabetes, and inflammation. Cancer alone is the second leading cause of death in the United States, accounting for perhaps 30% of all drug development effort globally.

The discovery of imatinib, an inhibitor that targets the Abelson tyrosine kinase (Abl) dysregulated in chronic myelogenous leukemia (CML) patients, was transformative in revealing the enormous therapeutic potential of selective kinase inhibitors, kindling hope that this remarkable success...
could be recapitulated for other cancers and diseases.\textsuperscript{7} While there are now 39 Food and Drug Administration-approved selective kinase small molecule inhibitors (as of January 16, 2018),\textsuperscript{8,9} these molecules were approved for targeting only 22 of ∼500 human kinases,\textsuperscript{10} with the vast majority developed to target just a handful of kinases.\textsuperscript{10} The discovery of therapeutically effective inhibitors for other kinases has proven to be remarkably challenging.

While these inhibitors have found success in the clinic, many patients cease to respond to treatment because of resistance caused by mutations in the targeted kinase,\textsuperscript{11} activation of downstream kinases,\textsuperscript{3} or relief of feedback inhibition in signaling pathways.\textsuperscript{12} These challenges have spurred the development of a new generation of inhibitors aimed at overcoming resistance,\textsuperscript{13,14} as well as mutant-specific inhibitors that target kinases bearing a missense mutation that confers resistance to an earlier-generation inhibitor.\textsuperscript{15} The ability to easily engineer and express mutant kinase domains of interest would be of enormous aid in the development of mutant-selective inhibitors, offering an advantage over current high-throughput assays,\textsuperscript{6,16–18} which typically include few clinically observed mutant kinases.

Probing human kinase biochemistry, biophysics, and structural biology in the laboratory is essential to making rapid progress in understanding kinase regulation, developing selective inhibitors, and studying the biophysical driving forces underlying mutational mechanisms of drug resistance. While human kinase expression in baculovirus-infected insect cells can achieve high success rates,\textsuperscript{19,20} it cannot compete in cost, convenience, or speed with bacterial expression. \textit{Escherichia coli} expression enables production of kinases without unwanted post-translational modifications, allowing for greater control of the system. A survey of 62 full-length nonreceptor human kinases found that >50\% express well in \textit{E. coli},\textsuperscript{19} but often expressing only the soluble kinase domains is sufficient, because these are the molecular targets of therapy for targeted kinase inhibitors and could be studied even for receptor-type kinases. While removal of regulatory domains can negatively impact expression and solubility, coexpression with phosphatase was shown to greatly enhance bacterial kinase expression in Src and Abl tyrosine kinases, presumably by ensuring that kinases remain in an unphosphorylated inactive form where they can cause minimal damage to cellular machinery.\textsuperscript{21}

The Protein Data Bank (PDB) now contains >100 human kinases that were expressed in bacteria, according to PDB header data. Many of these kinases were expressed and crystallized as part of the highly successful Structural Genomics Consortium (SGC) effort to increase structural coverage of the human kinome.\textsuperscript{22} Because bacterial expression is often complicated by the need to tailor construct boundaries, solubility-promoting tags, and expression and purification protocols individually for each protein expressed, we wondered whether a simple, universal, automatable expression and purification protocol could be used to identify tractable kinases, select construct boundaries, express a large number of human kinases and their mutant forms, and produce a convenient bacterial expression library to facilitate kinase research and selective inhibitor development. As a first step toward this goal, we developed a structural informatics pipeline to use available kinase structural data and associated metadata to select constructs from available human kinase libraries to clone into a standard set of vectors intended for phosphatase coexpression under a simple automatable expression and purification protocol. Using an expression screen for multiple

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**Figure 1.** Abl kinase domain construct expression screen that illustrates the high sensitivity to construct boundaries. (A) Abl kinase domain construct boundaries with the highest expression yields. Standard deviations of the yield are listed for control constructs for which six replicates were performed to give an indication of the uncertainty in experimental constructs. Secondary structure is indicated on the sequence. β-Sheets are colored blue, and α-helices are colored orange. (B) Heat map showing average yields for constructs (in micrograms per milliliter of culture) with detectable expression as a function of N- and C-terminal construct boundaries. (C) PDB entry 2E2B with the nine N-terminal construct boundary amino acids shown as yellow spheres (left) and PDB entry 4XEY with the nine C-terminal construct boundary amino acids shown as green spheres (right). Black arrows indicate residue numbers.

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From the Bench

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construct domain boundaries of Abl, we found that transferring construct boundaries from available structural data can produce constructs with useful expression levels, enabling simple identification of construct domain boundaries. We then completed an automated expression screen in Rosetta2 cells of 96 different kinases and found that 52 human kinase domains express with yields of $>2 \mu g/mL$ of culture. To investigate whether these kinases are properly folded and useful for biophysical experiments, we performed a fluorescence-based thermostability assay on the 14 highest-expressing kinases. Each well is marked with the Abl kinase domain construct residue boundaries (UniProt canonical isoform numbering). Bands for YopH164 phosphatase (50 kDa) and Abl kinase domain constructs (28–35 kDa) are labeled.

Briefly, the impact of construct boundary choice on Abl kinase domain expression was tested as follows (see Methods for full details). His10-TEV N-terminally tagged wild-type Abl constructs $^6$ were coexpressed with YopH phosphatase $^7$ in a 96-well format with control replicates distributed randomly throughout the plate. His-tagged protein constructs were recovered via a single nickel affinity chromatography step, and construct yields were quantified using microfluidic capillary electrophoresis following thermal denaturation. Expression yields are summarized in Figure 1A, and a synthetic gel image from the constructs with detectable expression is shown in Figure 2. Abl construct bands are present at sizes between 29 and 35 kDa (because of the variation in construct boundaries), and YopH phosphatase (which is not His-tagged but has substantial affinity for the nickel beads) is present in all samples at its expected size of 50 kDa. Strikingly, despite the fact that N-terminal and C-terminal construct boundaries varied over only 15–25 residues, only a small number of constructs produced detectable expression (Figure 1B). As highlighted in Figure 1C (left), the best N-terminal boundaries (residues 228–230) are located on a disordered strand distant from any secondary structure; N-terminal boundaries closer to the $\beta$-sheet of the N-lobe gave poor or no detectable expression (Figure 1B).

The best C-terminal construct boundaries (residues 511 and 512) occur in an $\alpha$-helix (Figure 1C, right). It is noteworthy that this $\alpha$-helix is not resolved in PDB entry 2E2B, $^{25}$ suggesting this structural element may only be weakly thermodynamically stable in the absence of additional domains. In previous work, this $\alpha$-helix was shown to undergo a dramatic conformational change that introduces a kink at residue 516, splitting the $\alpha$-helix into two. $^{26}$ This suggests a high potential for flexibility in this region.

Two of the control constructs (which differ in construct boundary by only one or two residues) were in the top six expressing constructs (Figure 1A) and were in fact within 60% of the maximum observed expression yield. From this, we concluded that transferring construct boundaries from existing kinase domain structural data would be sufficient to bias our constructs toward useful expression levels for a large-scale screen of multiple kinases.

**RESULTS**

**Construct Boundary Choice Impacts Abl Kinase Domain Expression.** To understand how alternative choices of expression construct boundaries can modulate bacterial expression of a human kinase domain, we performed an expression screen of 84 unique construct boundaries encompassing the kinase domain of the tyrosine protein kinase ABL1.

Three constructs known to express in bacteria were chosen from the literature and used as controls, spanning UniProt residues 229–500 (PDB entry 3CS9), $^{23}$ 229–512 (PDB entry 2G2H), $^{24}$ and 229–515 (PDB entry 2E2B); $^{25}$ 81 constructs were generated combinatorially by selecting nine different N-terminal boundaries spanning residues 228–243 and nine different C-terminal boundaries spanning residues 490–515, chosen to be near the start and end points for the control constructs, respectively (Figure 1A). Each of the three control constructs included six replicates to provide an estimate of the typical standard error in expression readout for the experimental constructs, which was found to be between 0.42 and 1.5 $\mu g/mL$ (Figure 1A, green constructs).

![Figure 2. Expression yields of Abl kinase domain constructs for all constructs with detectable expression. Synthetic gel image rendering generated from Caliper GX II microfluidic gel electrophoresis data following Ni affinity purification and thermal denaturation for all Abl constructs with detectable expression. Each well is marked with the Abl kinase domain construct residue boundaries (UniProt canonical isoform numbering). Bands for YopH164 phosphatase (50 kDa) and Abl kinase domain constructs (28-35 kDa) are labeled.](image-url)
truncated. This query returned a set of target sequences that were then matched to their relevant PDB constructs and filtered for expression system (as determined from PDB header EXPRESSION_SYSTEM records), discarding kinases that did not have any PDB entries with bacterial expression. As a final filtering step, the kinases were compared to three purchased kinase plasmid libraries (described in Methods), discarding kinases without a match. Construct boundaries were selected from PDB constructs and the SGC plasmid library, both of which have experimental evidence of E. coli expression, and subcloned from a plasmid in a purchased library (see Methods). Selecting the kinases and their constructs for this expression trial in this method rested on the basis of expected success: these specific kinase constructs were bacterially expressed and purified to a degree that a crystal structure could be determined. While expression protocols used to produce protein for crystallographic studies are often individually tailored, we considered these kinases to have a high likelihood of expressing in our semiautomated pipeline where the same protocol is utilized for all kinases. Statistics of the number of kinases obtained from the PDB mining procedure are shown in Figure 3A. Surprisingly, the most highly sampled family was the CAMK family, suggesting researchers may have found this family particularly amenable to bacterial expression. On the basis of the results of the previous experiment scanning Abl constructs for expression, we decided to use construct boundaries that were reported in the literature for each kinase. This process resulted in a set of 96 plasmid constructs distributed across kinase families (Figure 3B).

From these constructs, a set of 96 His10-TEV N-terminally tagged kinase domain constructs were generated, coexpressed with a phosphatase in E. coli, purified via nickel bead pull down, and quantified using microfluidic gel electrophoresis. The 96 kinases were coexpressed with either lambda phosphatase (for Ser/Thr kinases) or a truncated form of YopH phosphatase (for Tyr kinases).

Instead of eluting with imidazole, the purified kinase was cleaved off nickel beads by the addition of 10% TEV protease to minimize phosphatase contamination in the resulting eluate, allowing us to assess whether resulting yields would be sufficient (and sufficiently free of phosphatase) to permit activity assays. While the initial panel of 96 kinases was well-distributed among kinase families (Figure 3B), the most highly expressing kinases (yield of >12 μg of kinase/mL of culture) were not evenly distributed (Figure 3D). While many of the kinases chosen from the CMGC and CK1 families expressed well in our panel, nearly all of the kinases from the CAMK and AGC family express <12 μg of kinase/mL (Figure 3D); 52 kinases demonstrated a useful level of soluble protein expression, here defined as >2 μg/mL, naively expected to scale up to better than 2 mg/L of culture (Table 1). Some kinases (bold values in Table 1) demonstrated very high levels of expression, while others (plain type values in Table 1) would likely benefit from further rounds of construct boundary optimization or solubility tags to boost the level of soluble expression. The 17 most highly expressing kinases showed relatively high purity after elution, though we note that eluting via TEV site cleavage results in a quantity of TEV protease in the eluate (Figure 3C) but does not cause the elution of the His-tagged phosphatases that would hinder the ability to perform kinase activity assays. Further optimization of elution conditions may be required for optimizing kinase recovery via TEV cleavage.28−30

Figure 3. Kinomewide search for expressible kinases. (A) Number of PDB structures per kinase family, from the database built to select kinases for expression. (B) Distribution among families of candidate kinases in our expression screen. (C) Caliper GX II synthetic gel image rendering of the highest-expressing kinases, quantified using microfluidic capillary electrophoresis. (D) Kinome distribution of expression based on our 96-kinase screen. Dark green circles represent kinases with expression yields of >50 μg/mL of culture. Light green circles represent kinases with expression yields between 12 and 50 μg/mL. Yellow circles represent kinases with expression yields between 7 and 12 μg/mL. Orange circles represent kinases with any expression yield (even <2 μg/mL) up to 7 μg/mL. Image made with KinMap (http://www.kinhub.org/kinmap).
Table 1. Kinase Domain Constructs with Yields of >2 μg/mL of Culture for the 96-Kinase Expression Screen

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*Kinases are listed by UniProt designation and whether they were coexpressed with lambda or truncated YopH164 phosphatase. Yield (determined by Caliper GX II quantitation of the expected size band) reported in micrograms per milliliter of culture, where the total eluate volume was 120 μL from 900 μL of bacterial culture. Yields are shown in bold (yield > 12 μg/mL), italic (12 μg/mL > yield > 7 μg/mL), and plain type (yield < 7 μg/mL). Kinase domain constructs with yields that were undetectable or <2 μg/mL are not listed. † denotes that the second kinase domain of KS6A1_HUMAN was expressed; all other kinases were the first or only kinase domain occurring in the open reading frame. Construct boundaries are listed in UniProt residue numbering for the UniProt canonical isoform. An interactive table of expression yields and corresponding constructs is available at [http://choderalab.org/kinome-expression](http://choderalab.org/kinome-expression).
Constructs with expression yields of >2 \(\mu g/mL\) have been made available via Addgene (https://www.addgene.org/kits/chodera-kinase-domains).

High-Expressing Kinases Are Folded with a Well-Formed ATP Binding Site. To determine whether the expressed kinases were properly folded, we performed both a fluorescence-based thermostability assay (Figure 4) and a fluorescent ATP-competitive ligand binding measurement to quantify whether the ATP binding site was well-formed (Figure 5).

Fluorescence-Based Thermostability Assay. A fluorescence-based thermostability assay was performed with the hydrophobic dye SYPRO Orange to determine whether a strong two-state unfolding signal could be observed (see Methods). Also termed thermofluor or differential scanning fluorimetry (DSF), as the temperature is slowly increased, unfolded proteins will expose hydrophobic patches to which SYPRO Orange will bind, causing an increase in fluorescence.\(^{31-33}\) While the fluorescence of solvated SYPRO Orange is temperature-dependent, clear unfolding temperatures \((T_m)\) can often be identified from peaks in the first derivative of the observed fluorescence signal. Figure 4 shows the fluorescence (blue line), the absolute value of its derivative (red dashed line), and the unfolding temperature determined from the maximum absolute derivative (\(T_m\)) for the 14 kinases that were eluted to concentrations of >0.24 mg/mL after elution. SYPRO Orange fluorescence (solid blue line) was measured at 580 nm (half-bandwidth of 20 nm) after excitation at 465 nm (half-bandwidth of 25 nm) as the temperature was ramped (x-axis) in nickel buffer A [25 mM HEPES (pH 7.5), 5% glycerol, 400 mM NaCl, 20 mM imidazole, and 1 mM BME]. The temperature was held at 25 °C for 15 s before being increased to 95 °C with a ramp rate of 0.06 °C/s. Unfolding temperature \(T_m\) (black dashed line and inset) was determined from the maxima of the normalized first derivative of fluorescence (red dashed line). Fluorescence emission at 580 nm is shown on the left y-axis. To control for signals resulting from TEV protease contamination present at 0.01–0.03 mg/mL, TTK, a kinase with no detectable expression in our panel as determined via Caliper GX II quantitation, was included (panel 15).

Figure 4. Fluorescence-based thermostability assay that demonstrates many high-expressing kinases are well-folded. A fluorescence-based thermostability assay was performed on the 14 kinases shown to express above a minimum concentration of 0.24 mg/mL after elution. SYPRO Orange fluorescence (solid blue line) was measured at 580 nm (half-bandwidth of 20 nm) after excitation at 465 nm (half-bandwidth of 25 nm) as the temperature was ramped (x-axis) in nickel buffer A [25 mM HEPES (pH 7.5), 5% glycerol, 400 mM NaCl, 20 mM imidazole, and 1 mM BME]. The temperature was held at 25 °C for 15 s before being increased to 95 °C with a ramp rate of 0.06 °C/s. Unfolding temperature \(T_m\) (black dashed line and inset) was determined from the maxima of the normalized first derivative of fluorescence (red dashed line). Fluorescence emission at 580 nm is shown on the left y-axis. To control for signals resulting from TEV protease contamination present at 0.01–0.03 mg/mL, TTK, a kinase with no detectable expression in our panel as determined via Caliper GX II quantitation, was included (panel 15).
and 9), had two shallow inflection points in SYPRO fluorescence as a function of temperature. While STK3 does not have a strong peak above room temperature, titration with an ATP-competitive inhibitor suggests this kinase has a well-formed ATP binding site or folding can be induced by ligand binding (Figure 5, panel 10). As a control, a sample with no detectable kinase expression (TTK from our expression panel) was assayed (Figure 4, panel 9), which showed nearly no fluorescence signal.

**ATP-Competitive Inhibitor Binding Fluorescence Assay.**
To determine whether expressed kinases had well-folded ATP binding sites, we probed their ability to bind an ATP-competitive inhibitor. While a pan-kinase inhibitor such as staurosporine could be used as a fluorescent probe,35 the ATP-competitive inhibitor bosutinib shows a much stronger increase in fluorescence around 450−480 nm when bound to kinases with well-folded ATP binding sites.34,36 While excitation at 350 nm can be used, excitation at 280 nm results in a lower background, potentially because of fluorescent energy transfer between the kinase and ligand. Despite the weak affinity of bosutinib for many kinases, its aqueous solubility is sufficient to provide a quantitative assessment of ATP-competitive binding to many kinases at sufficiently high concentrations to function as a useful probe.34,36

Here, we utilized this approach as a qualitative probe for ATP-competitive ligand binding, because of uncertainty in the ligand concentration caused by significant evaporation over the course of the sequential titration experiment (see Methods for a more in-depth discussion); 33 of the kinases in our expression panel had sufficient yields to prepare 100 μL of 0.5 μM kinase assay solutions and were assessed for binding to bosutinib (Figure 5, panels 1−33), with a concentration-dependent increase in the intensity of the fluorescence signal above baseline qualitatively indicate the presence of a well-formed ATP binding site. Six of the lowest-expressing kinase constructs (Figure 5, panels 39−44) were prepared by diluting 20 μL to a reaction volume of 100 μL and assessed for bosutinib binding. Unexpectedly, these kinases also showed evidence of binding, suggesting this assay is able to detect a well-formed ATP binding site even for protein concentrations of <0.5 μM. To demonstrate that unfolded kinases do not demonstrate this increase in fluorescence over the ligand-only baseline, thermally denatured MK14 was included as a control next to folded MK14 from a large-scale expression preparation (Figure 5, panels 37 and 38), with thermally denatured MK14 exhibiting little difference from titrating ligand into buffer alone.

**Figure 5.** Fluorescence emission spectra as a function of the fluorescent ATP-competitive kinase inhibitor bosutinib demonstrate the presence of a well-formed ATP binding pocket. The ATP-competitive inhibitor bosutinib shows a strong increase in fluorescence centered around 450 nm when bound to kinases with well-folded ATP binding sites upon excitation at 280 nm.34 To assess whether the kinases from the high-throughput expression screen were well-folded, bosutinib was titrated in a 15-concentration series geometrically spanning 0.008−18.99 μM (colored lines; higher concentrations are shown in warmer colors) in 15 increments for 39 expressing kinases with the protein concentration adjusted to ~0.5 μM in a 100 μL assay volume. The eluted TEV protease contaminant concentration varies from 0.01 to 0.03 mg/mL in the assay volumes. The control MK14 and boiled MK14 (boiled for 10 min at 95 °C) were produced in a large-scale expression from the same plasmid that was used in the high-throughput expression protocol, and they were included as positive and negative controls for bosutinib binding to the ATP binding pocket. Fluorescence emission spectra (y-axis, bandwidth of 20 nm) were measured from 370 to 600 nm (x-axis) for excitation at 280 nm (bandwidth of 10 nm). For reference, the fluorescence of bosutinib titrated into buffer titration (panel 36) is shown in grayscale in each panel. Significant increases in the intensity of the fluorescence signal above baseline qualitatively indicate the presence of a well-formed ATP binding site.
Expressing Clinically Derived Src and Abl Missense Mutants. Next-generation sequencing has enabled generation of massive data sets rich with missense alterations in kinases observed directly in the clinic \(^{37-39}\) and has been particularly transformative in the field of oncology. To determine how well our human kinase domain panel supports the automated expression of clinically identified missense mutants for biophysical, biochemical, and structural characterization, we attempted to express 96 missense mutations mined from sequencing studies of cancer patients. The mutations were gathered using cBioPortal\(^40\) from publicly available sources and a large clinical tumor sequencing data set from the Memorial Sloan Kettering Cancer Center\(^38\) sequenced in the MSK-IMPACT panel.\(^41\)

Using our structural informatics pipeline, a database was built focusing on the kinases we found to be expressible in *E. coli*. To add the mutation data, we retrieved public data sets from cBioPortal\(^44,45\) along with annotations from Oncotator\(^46\) through their respective web service APIs. We then added mutations and annotations from the MSKCC data set\(^38\) by extracting the mutations from a local copy of the data set and retrieving annotations from Oncotator. The annotated mutations were filtered for mutations that occurred within the construct boundaries of our kinase domains. We found 63

![Figure 6. Expression yields for engineered clinically derived Src and Abl missense mutants. (A) All Abl and Src clinically identified mutants assessed in the expression screen are displayed as sticks. Mutants with expression yields of >2 μg/mL are colored green, while those with yields of <2 μg/mL are colored orange. Rendered structures are Abl (PDB entry 2E2B) and Src (PDB entry 4MXO).\(^36\) (B) Synthetic gel images showing Abl (top) or Src (bottom) expression, with wells labeled by missense mutation. The yield was determined by Caliper GX II quantitation of the expected size band and reported in micrograms per milliliter of culture, where the total eluate volume was 120 μL following nickel bead pull-down purification from a 900 μL bacterial culture. Residue mutations use numbering for the UniProt canonical isoform.](image-url)
unique clinical mutations appearing within our kinase domain construct boundaries for Abl and 61 for Src. We subsequently selected 48 mutants for Abl and 46 for Src to express, aiming for a panel of mutants distributed throughout the kinase domain (Figure 6A), with wild-type sequences included as controls. Mutations were introduced using site-directed mutagenesis and assayed for expression yields (Figure 6B). Those with yields of >2 μg of kinase/mL of culture are listed in Table 2.

High-expressing mutants appear to be distributed relatively uniformly throughout the kinase domain (Figure 6A). While the vast majority of the Src mutants expressed at a usable level, many of the Abl mutants expressed below the 2 μg/mL threshold. This can primarily be attributed to the low level of expression for the wild-type Abl construct (Table 1). In instances in which kinase activity is not required, the yield could be increased via the introduction of inactivating mutations23 or further tailoring of expression and purification protocols.

Table 2. Expression Yields for Engineered Clinical Missense Mutants of Src and Abl Kinase Domains with Yields of >2 μg/mL of Culture

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“Src and Abl kinase domain constructs with engineered clinical mutations with expression yields of >2 μg/mL of culture are listed, sorted by yield. The yield was determined by Caliper GX II quantitation of the expected size band and reported in micrograms per milliliter of culture, where the total eluate volume was 80 μL purified from a 900 μL bacterial culture. Wild-type (WT) controls for both Src and Abl (here, a single well for each) are shown as the first entry for each gene. The UniProt amino acid sequence numbering of the primary isoform. MutationAssessor Score,42,43 which predicts functional impact via conservation.

■ METHODS

Semiautomated Selection of Kinase Construct Sequences for E. coli Expression. Selection of Human Protein Kinase Domain Targets. Human protein kinases were selected by querying the UniProt API (query date of May 30, 2014) for any human protein with a domain containing the string “protein kinase” and that was manually annotated and reviewed (i.e., a Swiss-Prot entry). The query string used was taxonomy:”Homo sapiens” (Human) [9606]” AND domain:”protein kinase” AND reviewed:yes.

Data were returned by the UniProt API in XML format and contained protein sequences and relevant PDB structures, along with many other types of genomic and functional information. To select active protein kinase domains, the UniProt domain annotations were searched using the regular expression “Protein kinase; inactive,” which excludes certain domains annotated “Protein kinase; truncated” and “Protein kinase; inactive.” Sequences for the selected domains, derived from the canonical isoform as determined by UniProt, were then stored.

Matching Target Sequences with Relevant PDB Constructs. Each target kinase gene was matched with the
homologues in any other species, if present, and all UniProt data were downloaded. These data included a list of PDB structures that contain the protein and their sequence spans in the coordinates of the UniProt canonical isoform. PDB structures that did not include the protein kinase domain or had more than 30 residues at each end truncated were filtered out. PDB coordinate files were then downloaded for each remaining PDB entry. The coordinate files contain various metadata, including the EXPRESSION_SYSTEM annotation, which was used to filter PDB entries for those which include the phrase "ESCHERICHIA COLI." The majority of PDB entries returned had an EXPRESSION_SYSTEM tag of "ESCHERICHIA COLI," while a small number had "ESCHERICHIA COLI BL21" or "ESCHERICHIA COLI BL21(DE3)."

The PDB coordinate files also contain SEQRES records, which should contain the protein sequence used in the crystallography or NMR experiment. According to PDB-101 (http://pdb101.rcsb.org/learn/guide-to-understanding-pdb-data/primary-sequences-and-the-pdb-format), the SEQRES should include the "sequence of each chain of linear, covalently linked standard or modified amino acids or nucleotides. It may also include other residues that are linked to the standard backbone in the polymer." However, we found that these records are very often misannotated, instead representing only the crystallographically resolved residues. Because expression levels can be greatly affected by insertions or deletions of only one or a few residues at either terminus, it is important to know the full experimental sequence. To measure the authenticity of a given SEQRES record, we developed a simple metric by hypothesizing that most crystal structures would likely have at least one or more unresolved residues at one or both termini and that the presence of an expression tag, which is typically not crystallographically resolved, would indicate an authentic SEQRES record. To achieve this, unresolved residues were first defined by comparing the SEQRES sequence to the resolved sequence, using the SIFTS service to determine which residues were not present in the canonical isoform sequence. Regular expression pattern matching was used to detect common expression tags at the N- or C-termini. Sequences with a detected expression tag were given a score of 2, while those with any unresolved sequence at the termini were given a score of 1; the remainder were given a score of 0. These data were stored to allow for subsequent selection of PDB constructs based on likely authenticity in later steps. The number of residues extraneous to the target kinase domain and the number of residue conflicts with the UniProt canonical isoform within that domain span were also stored for each PDB sequence.

**Plasmid Libraries.** As a source of kinase DNA sequences for subcloning, we purchased three kinase plasmid libraries: the Addgene Human Kinase ORF kit, a kinase library from the Structural Genomics Consortium (SGC), Oxford (http://www.thesgc.org), and a kinase library from the PlasmID Repository maintained by the Dana-Farber/Harvard Cancer Center. Annotated data for the kinases in each library were used to match them to the human protein kinases selected for this project. The plasmid open reading frames (ORFs) were translated into protein sequences and aligned against the target kinase domain sequences from UniProt. Also calculated were the numbers of extraneous protein residues in the ORF, relative to the target kinase domain sequence, and the numbers of residue conflicts with the UniProt sequence. Our aim was to subclone the chosen sequence constructs from these library plasmids into our expression plasmids.

**Selection of Sequence Constructs for Expression.** Of the kinase domain targets selected from UniProt, we filtered out those with no matching plasmids in our available plasmid libraries or no suitable PDB construct sequences. For this purpose, a suitable PDB construct sequence was defined as any with an authenticity score greater than zero (see above). Library plasmid sequences and PDB constructs were aligned against each UniProt target domain sequence, and various approaches were considered for selecting the construct boundaries to use for each target and the library plasmid from which to subclone it. Candidate construct boundaries were drawn from two sources, PDB constructs and the SGC plasmid library, which has been successfully tested for E. coli expression.

For most of the kinase domain targets, multiple candidate construct boundaries were available. To select the most appropriate construct boundaries, we sorted them first by authenticity score, then by the number of conflicts relative to the UniProt domain sequence, and then by the number of residues extraneous to the UniProt domain sequence span. The top-ranked construct was then chosen. In cases in which multiple library plasmids were available, these were sorted first by the number of conflicts relative to the UniProt domain sequence and then by the number of residues extraneous to the UniProt domain sequence span, and the top-ranked plasmid was chosen. This process resulted in a set of 96 kinase domain constructs, which (by serendipity) matched the 96-well plate format we planned to use for parallel expression testing. We selected these constructs for expression testing.

An interactive table of the selected plasmids, constructs, and aligned PDB files can be viewed at http://choderalab.org/kinome-expression.

**Automation of the Construct Selection Process.** While much of this process was performed programmatically, many steps required manual supervision and intervention to correct for exceptional cases. While these exceptions were encoded programmatically as overrides to ensure the scheme could be reproduced from existing data, we hope to eventually develop a fully automated software package for the selection of expression construct sequences for a given protein family, but this was not possible within the scope of this work.

**Mutagenesis Protocol.** Point mutations were introduced with a single-primer QuickChange reaction. Primers were designed to anneal at 55 °C both upstream and downstream of the point mutation, with a total length of approximately 40 bases. At the codon to be modified, the fewest possible number of bases was changed. The plasmid template (160 ng) was mixed with 1 μM primer in 1X PfuUltra reaction buffer, with 0.8 mM dNTPs (0.2 mM each) and 1 unit of PfuUltra high-fidelity DNA polymerase (Agilent), in a total volume of 20 μL. Thermocycler settings were 2 min at 95 °C, followed by 18 cycles of 20 s at 95 °C, 1 min at 53 °C, 12 min at 68 °C (2 min/kb), and then 1 min at 68 °C. After the mixture had cooled to room temperature, 4 μL of the polymerase chain reaction (PCR) mixture was added to 16 μL of CutSmart Buffer (NEB) containing 10 units of DpnI (NEB). After incubation for 2.5 h at 37 °C, 6 μL of this mixture was used to directly transform XLI-Blue chemically competent cells (Agilent) according to the manufacturer’s protocol. Transformants were picked for plasmid mini-preps, and the presence of the point mutations was confirmed by sequencing.
Expression Testing. For each target, the selected construct sequence was subcloned from the selected DNA plasmid. Expression testing was performed at the QB3MacroLab (http://qb3.berkeley.edu/macrolab/), a core facility offering automated gene cloning and recombinant protein expression and purification services.

Each kinase domain was tagged with an N-terminal His10-TEV and coexpressed with either the truncated YopH164 for Tyr kinases or lambda phosphatase for Ser/Thr kinases. All construct sequences were cloned into the 2BT10 plasmid, an AMP resistant ColE1 plasmid with a T7 promoter, using ligation-independent cloning (LIC). The inserts were generated by PCR using the LICv1 forward (TACTTCCAATCCAATGCA) and reverse (TTATCCACTTCCAATGTTATTA) tags on the primers. Gel-purified PCR products were treated via LIC with dCTP. The LIC-treated plasmid and insert were mixed together and transformed into XL1-Blue cells for plasmid preps.

Expression was performed in Rosetta2 cells (Novagen) grown with MagicMedia (Invitrogen autoinducing medium), 100 μg/mL carbenicillin, and 100 μg/mL spectinomycin. Single colonies of transformants were cultivated with 900 μL of MagicMedia in a gas permeable sealed 96-well block. The cultures were incubated at 37 °C for 4 h and then at 16 °C for 40 h while being shaken. Next, cells were centrifuged, and the pellets were frozen at −80 °C overnight. Cells were lysed on a rotating platform at room temperature for 1 h using 700 μL of SoluLye (Genlantis) supplemented with 400 mM NaCl, 20 mM imidazole, 1 μg/mL pepstatin, 1 μg/mL leupeptin, and 0.5 mM phenylmethylsulfonyl fluoride.

For protein purification, 500 μL of the soluble lysate was added to a 25 μL Protopo Ni-NTA (Machery-Nagel) agarose resin in a 96-well filter plate. Nickel buffer A [25 mM HEPES (pH 7.5), 5% glycerol, 400 mM NaCl, 20 mM imidazole, and 1 mM BME] was added, and the plate was shaken for 30 min at room temperature. The resin was washed with 2 mL of nickel buffer A. For the 96-kine expression experiment, target proteins were eluted by a 2 h incubation at room temperature with 10 μg of TEV protease in 80 μL of nickel buffer A per well and a subsequent wash with 40 μL of nickel buffer A to maximize protein release. Nickel buffer B [25 mM HEPES (pH 7.5), 5% glycerol, 400 mM NaCl, 400 mM imidazole, and 1 mM BME] was used to elute TEV resistant material remaining on the resin. Untagged protein eluted with TEV protease was run on a LabChip GX II Microfluidic system to analyze the major protein species present.

For the clinical mutant and Abi1 construct boundary expression experiments, target proteins were washed three times with nickel buffer A prior to elution in 80 μL of nickel buffer B. The eluted protein was run on a LabChip GX II Microfluidic system to analyze with major protein species present.

Fluorescence-Based Thermostability Assay. To assess whether the highly expressed wild-type kinase constructs are folded, a thermofluor thermostability assay31–33 was performed for kinase constructs that have a minimum protein concentration of 0.24 mg/mL in the eluate. After dilution of 9 μL of eluate with 1 μL of dye, the effective assay concentration is ≥0.216 mg/mL in a 10 μL assay volume. Previous optimization efforts in the lab determined that 0.20 mg/mL was the lower limit of well-defined Tm detection. This minimum concentration also ensured that the kinase was present at a concentration roughly an order of magnitude higher than that of the contaminating TEV protease.

Kinase expression panel eluates, which were kept in a 96-well deep well plate frozen at −80 °C for 2 years prior to the thermal stability assay, were thawed in an ice-water bath for 30 min. Nine microliters of each kinase eluate was added to a 384-well PCR plate (4itude-0381). The 100X SYPRO Orange dye solution was prepared from a 5000X DMSO solution of SYPRO Orange Protein Gel Stain (Life Technologies, reference no. S6650, lot no. 1790705) by dilution in distilled water. In initial experiments, the SYPRO Orange dye solution was diluted in kinase binding assay buffer [20 mM Tris and 0.5 mM TCEP (pH 8)], which caused the dye to precipitate out of the solution. Particles in the dye solution were pelleted by tabletop centrifugation (2 min, 5000 RCF), and the solution was kept covered with aluminum foil in the dark to prevent photodamage. One microliter of the 100X dye solution was added to each kinase eluate sample in a 384-well PCR plate. The plate was sealed with Axygen UC-500 Ultra Clear Pressure Sensitive sealing film. To remove any air bubbles, the sample plate was centrifuged for 30 s at 250g using a Bionex HiG4 centrifuge. Sample mixing was performed by orbital shaking with InHeco shakers for 2 min at 1000 rpm.

A thermofluor melt was performed using a LightCycler 480 (Roche) qPCR instrument using an excitation filter at 465 nm (half-bandwidth of 25 nm) and an emission filter at 580 nm (half-bandwidth of 20 nm). LightCycler 480 Software version 1.5.1 was used to operate the instrument and analyze the results. The temperature was held at 25 °C for 15 s before being increased to 95 °C at a ramp rate of 0.06 °C/s. During the temperature ramp, 10 fluorescence acquisitions/°C were recorded in the dynamic integration time mode, with a melt factor of 1, a quant factor of 10, and a maximum integration time of 2 s. Thermal protein denaturation causes hydrophobic patches of protein to be exposed, which SYPRO Orange dye can bind. Binding of SYPRO Orange dye is detected as an increase in fluorescence at 580 nm. The presence of a clear thermal denaturation peak in the absolute value of the derivative of the fluorescence as a function of temperature serves as an indication that the proteins were well-folded. The observed fluorescence was plotted as a function of temperature, and a melting temperature Tm was determined as the maximum of the absolute value of its first derivative.

ATP-Competitive Inhibitor Binding Fluorescence Assay. To determine whether the expressed kinases had a well-folded ATP binding site, we assessed whether the eluted kinase was capable of binding the ATP-competitive small molecule kinase inhibitor bosutinib. We designed fluorescence-based binding assays following earlier work reporting that this quinoline inhibitor undergoes a strong increase in fluorescence upon binding (even weakly) to kinase ATP binding sites.34 Via titration of the ligand close to the solubility limit, even weak binding to the ATP binding site can be detected by observing emission increases around 450 nm during excitation at 280 nm. For 33 of the kinases in our expression panel, 0.5 μM kinase solutions from kinase expression panel eluates were prepared in kinase binding assay buffer [20 mM Tris and 0.5 mM TCEP (pH 8)] for a final volume of 100 μL in a black 96-well vision plate (4itude-0223). Six low-expressing kinases (Figure S6 panels 39–44) were prepared by diluting 20 μL of the eluate in kinase binding assay buffer [20 mM Tris and 0.5 mM TCEP (pH 8)] until the final volume reached 100 μL, for final concentrations of <0.5 μM. The plate was shaken for 2 min
clockwise and 2 min counterclockwise by orbital shaking with
Inheco shakers at 2000 rpm and centrifuged for 30 s at 1000g
using a Bionex HiG4 centrifuge. Fluorescence emission spectra
were measured from 370 to 600 nm (20 nm bandwidth) in 5
nm steps using 280 nm excitation (10 nm bandwidth) from
both the top and bottom of the well using a Tecan Infinite
M1000 PRO instrument.

Bosutinib free base (LC Laboratories, catalog no. B-1788, lot
no. BSB-103, molecular weight of 530.45 Da) was dispensed
directly from a roughly 10 mM DMSO stock solution to the
assay solution using a Tecan HP D300 Digital Dispenser. The
10 mM DMSO stock solution was prepared gravimetrically
using an automated balance (Mettler Toledo Balance XPE205
with LabX Laboratory Software) by dispensing 39.02 mg of
solid bosutinib powder stored under nitrogen gas at 25 °C into
8.0499 g of DMSO (Alfa Aesar, catalog no. 42780, log no.
Y2SB604, density of 1.1004 g/mL at ambient temperature)
that is kept dry under argon gas at 25 °C. To minimize
atmospheric water absorption due to the hygroscopic nature of
DMSO, the 10 mM stock solution was pipetted into wells of a
96-well stock plate by an automated liquid handling device
(Tecan EVO 200 with air LiHa) and sealed with foil seal
(PlateLoc). The ligand was dispensed to the assay plate with
HP D300 (using aliquots of stock solution pipetted from a
freshly pierced stock plate well) targeting a roughly geometri-
cal shape (PlateLoc). The ligand was dispensed to the assay plate with
HP D300 (using aliquots of stock solution pipetted from a
freshly pierced stock plate well) targeting a roughly geometri-
cally increasing series of ligand concentrations in each well to
achieve the following total ligand concentrations after each dispensing:
0.008, 0.013, 0.023, 0.038, 0.064, 0.109, 0.183,
0.308, 0.519, 0.875, 1.474, 3.174, 6.037, 10.862, and 18.991
μM. The plate was shaken with an HP D300 instrument for 10
s after the use of each dispensehead. After each titration, the
plate was shaken with Inheco shakers (2 min clockwise and
counterclockwise, 2000 rpm, orbital shaking) and centrifuged
(30 s at 1000g) using a Bionex HiG4 centrifuge. Fluorescence
spectra from 370 to 600 nm (bandwidth of 20 nm) in 5 nm steps using 280 nm excitation (bandwidth of 10 nm) were read
from both the top and the bottom of the well using a Tecan
Infinite M1000 PRO instrument. In total, the experiment took
17.5 h to complete because of the time-consuming spectral
read after each dispensing, likely resulting in significant
evaporation from some wells during the experiment.

ATP-competitive binding was analyzed qualitatively for each
kinase by plotting the fluorescence spectra as a function of
concentration to detect concentration-dependent increases in
fluorescence. As a control for background ligand fluorescence
independent of protein binding, fluorescence spectra of three
replicates of ligand into buffer titrations were plotted. As a
positive control, MK14 produced by a validated large-scale
expression protocol (see the Supplementary Methods) from
the same plasmid used in the high-throughput protocol was
included. To control for nonspecific binding to unfolded protein, we included boiled MK14 (prepared from the large-
scale expression of MK14 by boiling at 95 °C for 10 min). A
concentration-dependent increase in fluorescence was inter-
preted as evidence that the ATP binding site of the kinase was
well folded and allowed for bosutinib binding. Because of the
length of the experiment, it is possible that evaporation
reduced the well volume below 100 μL and potentially caused
bosutinib to reach higher than expected concentrations. This
creates uncertainty for data points, as bosutinib may be at
either a higher-than-expected concentration (due to evapo-
ratation) or a lower-than-expected concentration (due to
potential precipitation caused by smaller well volumes). For
this reason, we have interpreted the experiment as qualitative
evidence of binding, instead of quantitative. Bosutinib binding
is an indication of proper folding of the ATP binding pocket of
these recombinantly expressed kinase constructs.

### DISCUSSION

We have demonstrated that a simple, uniform, automatable protocol is able to achieve useful bacterial expression yields for
a variety of kinase domain constructs. While yields could likely
be further improved by a variety of methods—such as the
addition of solubility-promoting tags, construct domain
boundary and codon optimization, or mutations to improve
the solubility or ablate catalytic activity—the simplicity of this
approach suggests widespread utility of automated bacterial
expression for biophysical, biochemical, and structural biology
work for the further study of human kinase domains.

Our expression test of 81 different construct boundaries of the Abl kinase domain demonstrated a surprising sensitivity of
expression yields to the precise choice of boundary. This
sensitivity may be related to where the construct is truncated
with respect to the secondary structure of the protein, as
disrupting the secondary structure could cause the protein to
improperly fold, leading to a low soluble protein yield even
when the total expression level is high. It is noteworthy that the
highest-expressing C-terminal boundaries for Abl were residues
511 and 512. These residues fall in regulatory α-helix e8.26
This helix has been shown to undergo a dramatic conforma-
tional change upon binding to the myristoylated N-terminal
cap, which introduces a sharp “kink” in residues 516–519.
These residues may lead to higher levels of soluble expression
by truncating an secondary structural element that is unusually
flexible. Indeed, this helix is not resolved in some X-ray
structures (PDB entry 2E2B), further suggesting that this helix
is less thermodynamically stable than expected. Control
replicates of three constructs indicate good repeatability of
expression yields in the high-throughput format. This screen
suggests that optimization of construct boundaries could
potentially further greatly increase yields of poorly expressing
kinase domains. Codon optimization for bacterial expression
could also increase the level of expression for kinase domains
with low yields due to codon bias, as could coexpression with
chaperones.50

For those kinases that did express, a fluorescence-based
thermostability assay indicated that many of the highest-
expressing kinases are well-folded. An ATP-competitive
inhibitor binding fluorescent assay provides qualitative
evidence that the 39 kinases that had sufficiently high
expression levels to be assayed have a well-formed ATP
binding site capable of binding bosutinib, a small molecule
ATP-competitive kinase inhibitor. Taken together, these two
experiments demonstrate that our expression protocol
produces folded kinases with utility for biophysical experi-
ments and drug design.

The tolerance of these bacterial constructs to many
engineered clinical missense mutations suggests a promising
route to the high-throughput biophysical characterization of
the effect of clinical mutants on anticancer therapeutics.
Mutations that did not express well may destabilize the protein
or may increase the specificity of the kinase. A higher
specific activity would require more phosphatase activity,
wasting ATP to prevent high levels of phosphorylation that
have been hypothesized to cause difficulty expressing kinases
without a coexpressed phosphatase in bacteria.21 Mutations
that are destabilizing may show improved expression if coexpressed with more elaborate chaperones such as GroEL and Trigger factor.\textsuperscript{50} Mutations that increase the specific activity of the kinase might also express better when combined with an inactivating mutation.

High-throughput automated kinase expression could be combined with enzymatic or biophysical techniques for characterizing the potency of a variety of clinical kinase inhibitors to assess which mutations confer resistance or sensitivity. While the process of engineering, expressing, purifying, and assaying mutants currently takes approximately 2 weeks, it is possible that new techniques for cell-free bacterial expression\textsuperscript{51,52} may reduce this time to a matter of days or hours in a manner that might be compatible with clinical time frames to impact therapeutic decision making.

We hope that other laboratories find these resources useful in their own work.

\section*{ASSOCIATED CONTENT}

\subsection*{Supporting Information}

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.biochem.7b01081.

Method for large-scale expression and purification of MK14\textsuperscript{(PDF)} spreadsheet containing the full expression results for all ABL1 constructs, wild-type kinases, and ABL1 and SRC mutants\textsuperscript{(XLSX)}

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Notes

The authors declare the following competing financial interest(s): J.D.C. is a member of the Scientific Advisory Board for Schrodinger, LLC.

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\section*{ADDITIONAL NOTES}

These targets are, currently, Abl, DDR1, EGFR, HER2, VGFRI/2/3, Alk, Met, BRAF, JAK1/2/3, Btk, Pi3K, CDK4, CDK6, MEK, ROS1, FLT3, IGFIIR, Ret, Kit, Axl, TrkB, and mTOR.

\textsuperscript{5}The parent plasmid is a pET His10 TEV LIC cloning vector and is available on Addgene (plasmid 78173).

\textsuperscript{6}Yoph164 phosphatase, engineered to minimize intrinsic affinity for nickel purification resin by the QB3MacroLab based on parent plasmid pCDFDuet1-YOPH, a gift from the Kuriyan lab.

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