

Data Mining Crystallization Databases: Knowledge-Based Approaches to Optimize Protein Crystal Screens

Matthew S. Kimber,¹ François Vallee,¹ Simon Houston,¹ Alexander Nečakov,¹ Tatiana Skarina,² Elena Evdokimova,² Steven Beasley,² Dinesh Christendat,² Alexei Savchenko,² Cheryl H. Arrowsmith,^{1,2} Masoud Vedadi,¹ Mark Gerstein,³ and Aled M. Edwards^{1,2*}

¹Affinium Pharmaceuticals Inc., Toronto, Ontario, Canada

²Division of Molecular and Structural Biology, Ontario Cancer Institute and Department of Medical Biophysics, University of Toronto, Toronto, Ontario, Canada

³Molecular Biophysics and Biochemistry, Yale University, New Haven, Connecticut

ABSTRACT Protein crystallization is a major bottleneck in protein X-ray crystallography, the workhorse of most structural proteomics projects. Because the principles that govern protein crystallization are too poorly understood to allow them to be used in a strongly predictive sense, the most common crystallization strategy entails screening a wide variety of solution conditions to identify the small subset that will support crystal nucleation and growth. We tested the hypothesis that more efficient crystallization strategies could be formulated by extracting useful patterns and correlations from the large data sets of crystallization trials created in structural proteomics projects. A database of crystallization conditions was constructed for 755 different proteins purified and crystallized under uniform conditions. Forty-five percent of the proteins formed crystals. Data mining identified the conditions that crystallize the most proteins, revealed that many conditions are highly correlated in their behavior, and showed that the crystallization success rate is markedly dependent on the organism from which proteins derive. Of the proteins that crystallized in a 48-condition experiment, 60% could be crystallized in as few as 6 conditions and 94% in 24 conditions. Consideration of the full range of information coming from crystal screening trials allows one to design screens that are maximally productive while consuming minimal resources, and also suggests further useful conditions for extending existing screens. *Proteins* 2003;51:562–568.

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INTRODUCTION

The ultimate goal of structural proteomics is to obtain, through experimental or computational methods, 3D structural models for every protein in nature. Driving this ambitious program is the expectation that structural information will provide functional insights for many of the proteins predicted by genome-sequencing efforts that cannot be ascribed a function using current sequence homology-based approaches. The challenges in structural

proteomics are significant; it has been estimated that some 16,000 structures will have to be determined using experimental approaches to obtain reasonable coverage of fold space.¹ Recent advances in X-ray crystallography methodology and associated technologies have made it possible, at least in ideal cases, to go from data collection to a refined structure in a matter of hours^{2–6}; however, actually growing a diffraction-quality crystal is far more time and resource intensive.

The purpose of protein crystallization trials is to efficiently find useful lead conditions from which crystal size and morphology can be optimized. Commonly, one explores a wide range of different solutions in which the salt concentration and type, pH, additive type, temperature, and precipitant type and concentration are varied. The precipitant is usually a long-chain polymer [poly-ethylene glycol (PEG), jeffamine], an inorganic or organic salt, or some small organic molecule (MPD, isopropanol, ethanol).²

There are two general approaches to the design of crystallization screens. One strategy aims to blanket potentially useful crystallization space with screens of a few hundreds to over a thousand conditions (see, e.g., the JBScreen at www.jenabioscience.com/jbscreen.html). The other strategy is to use smaller, more efficient screens based on previously successful conditions.^{3–8} The most widely used variant of this second strategy, developed by Jancarik and Kim, is based on an incomplete factorial approach, which explores a range of conditions biased toward previously successful crystallization conditions.⁸ The popularity of this screening strategy can be ascribed to many reasons, including its economy (1–2 mg of protein are needed), ease of use, manageable size, and convenience (it is sold in preformulated kit form by various companies). As originally formulated, this screen was intended to be modified reiteratively as the experiences of users were incorporated.³ In practice, however, this has not happened. Partly this is due to a lack of a clear metric by which

*Correspondence to: Aled M. Edwards, Affinium Pharmaceuticals Inc., 10th floor, South Tower, 100 University Avenue, Toronto, Ontario M5J 1V6, Canada. E-mail: aled.edwards@utoronto.ca

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TABLE I. Breakdown of Results by Source Organism

Organism	Number of proteins screened	Number of proteins crystallized	Mean hits per protein crystallized ^a	% success
<i>S. aureus</i>	372	142	4.2	38.2
<i>H. pylori</i>	128	47	6.4	36.7
<i>E. coli</i>	116	72	4.8	62.1
<i>M. thermoautotrophicum</i>	95	41	4.8	43.2
<i>Th. maritima</i>	34	23	3.5	67.6
<i>P. aeruginosa</i>	21	13	6.0	61.9
Overall	755	338	4.7	44.7

^aThe average number of conditions under which crystals were obtained, considering only those samples for which at least one crystal was obtained.

to decide which of a set of potential screening conditions is better, but also, more fundamentally, a lack of sufficient, standardized experimental data by which to evaluate a screen. While the optimal conditions for crystallizing a particular protein crystal form are often collected and archived in a database,⁹ the detailed results of each screen, including partial successes or failures, are not. As a result, while individual crystallization conditions can perhaps be shown to be more or less successful, researchers have tended to supplement the common factorial screens with additional screens rather than attempting to systematically optimize any given one. This may be a reasonable strategy when trying to crystallize one or two proteins, in which the effort and expense involved in making the additional protein, setting up the screen, and evaluating the results is manageable. However, in the context of structural proteomics this extra effort and expense is multiplied over hundreds or even thousands of proteins, and thus the desirability of screens that minimize the number of experiments while maximizing the probability of success becomes far more pronounced. Here, by collating data collected for 755 proteins from 6 organisms we show that it is possible to use the information gleaned from previous screens to improve screening strategies using objective empirical criteria.

METHODS

Proteins predicted not to have membrane-spanning domains from six organisms—the prokaryotes *Staphylococcus aureus*, *Escherichia coli* K12, *Pseudomonas aeruginosa*, *Helicobacter pylori*, and *Thermotoga maritima* and the archaeote *Methanobacterium thermoautotrophicum*—were amplified by PCR, cloned into *E. coli* expression vectors, overexpressed, and purified using His₆ technology, as described elsewhere (see [10] for a general overview and [11, 12] for typical examples of procedures). Proteins were typically stored at 4°C, in 20 mM HEPES, pH 7.5, and 500 mM NaCl. The solutions for the initial screen were purchased from Hampton Research. Proteins were screened in 24-well Lindbro plates, using a 2-μl + 2-μl drop size and 700 μl in the well. For some proteins both the his-tagged and non-his-tagged sample were both screened, and these were scored as separate samples. Samples also included separate domains of multidomain proteins. For most samples, two to four protein concentrations, typically

ranging from 5–40 mg ml⁻¹, were screened in parallel; in almost all cases this included at least one experiment in the 10- to 15-mg/ml range. The data for these multiple experiments were pooled. All crystallization experiments were performed at ambient temperature (approximately 293 K). In total, over 35,000 experiments were performed.

Screening results were scored by eye after approximately 1 day, 1 week, 1 month, and 3 months. For proteins where screening multiple samples at different protein concentrations yielded different outcomes, only the most favorable outcome was scored and reported. To minimize subjectivity in scoring, results for each experiment were reduced to one of three assessments—clear, precipitate, or crystalline. Samples were required to have at least two conditions in which they are soluble and two where they are not and no more than five conditions for which data was missing (if, e.g., the condition was not set or the drop dried out before it could be scored). These criteria reject 6 *E. coli* proteins, 3 *T. maritima* proteins, 6 *M. thermoautotrophicum* proteins, and 55 *S. aureus* proteins. Minimal screen 6 was derived by sequentially searching all combinations of conditions (48!/6! 42! ≈ 1.2 × 10⁷ combinations) for the one that crystallized the most proteins. Minimal screen 12 was derived by using minimal screen 6 as a seed and searching all combinations of the remaining conditions for the six that best complemented minimal screen 6. Minimal screen 24 was found by repeating this condition twice more. Although this procedure is not guaranteed to find the globally optimal subsets, this limitation is likely far less serious than the one imposed by the limited amount of data available. Clustering was performed in ClustalX using pairwise identity scores once screening results had been “encoded” as amino acid sequences.¹³ Cladogram was produced in Phylodraw.¹⁴

RESULTS AND DISCUSSION

A total of 755 protein samples from *T. maritima*, *E. coli*, *M. thermoautotrophicum*, *S. aureus*, *P. aeruginosa*, and *H. pylori* were screened against conditions 1–48 of the Janarik and Kim screen. The success rates for different genomes (Table I) ranged from 67.6% (for *T. maritima*, albeit with the smallest sample size) to 36.7% in the case of *H. pylori*. The differences may reflect differences in the intrinsic properties of proteins from different organisms, perhaps influenced by the intracellular environment of the

TABLE II. Overview of Results Produced by the Jancarik and Kim Screen[†]

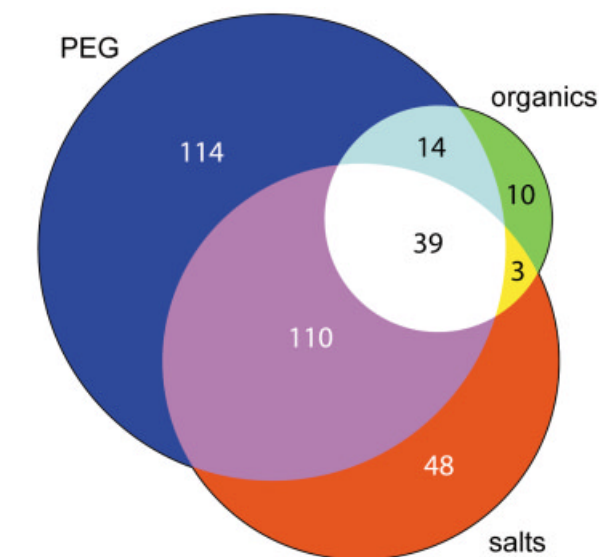
Jancarik–Kim no.	Components of screening condition	pH ^a	Total clear	Total precipitate	Total crystals	Only crystal ^b
1	30% MPD Na Acetate pH 4.6 0.02 M CaCl ₂	5.06	197	540	17	1
2	0.4 M K ₂ Na Tartrate	7.27	609	138	8	0
3	0.4 M NH ₄ Phosphate	4.24	264	450	12	0
4	2.0 M NH ₄ Sulfate Tris.HCl pH 8.5	8.31	208	494	50	3
5	30% MPD Na Hepes pH 7.5 0.2 M Na Citrate	7.44	346	396	11	0
6	30% PEG 4000 Tris.HCl pH 8.5 0.2 M MgCl ₂	8.70	84	601	65	4
7	1.4 M Na Acetate Na Cacodylate pH 6.5	6.83	513	211	27	0
8	30% Isopropanol Na Cacodylate pH 6.5 0.2 M Na Citrate	7.06	201	544	9	1
9	30% PEG 4000 Na Citrate pH 5.6 0.2 M NH ₄ Acetate	6.54	108	568	76	1
10	30% PEG 4000 Na Acetate pH 4.6 0.2 M NH ₄ Acetate	5.82	70	632	49	4
11	1.0 M NH ₄ Phosphate Na Citrate pH 5.6	4.89	309	404	15	0
12	30% Isopropanol Na Hepes pH 7.5 0.2 M MgCl ₂	7.29	177	560	16	1
13	30% PEG 400 Tris.HCl pH 8.5 0.2 M Na Citrate	8.84	551	193	10	2
14	28% PEG 400 Na Hepes pH 7.5 0.2 M CaCl ₂	7.32	212	517	25	2
15	30% PEG 8000 Na Cacodylate pH 6.5 0.2 M NH ₄ Sulfate	6.68	123	568	60	1
16	1.5 M Li Sulfate Na Hepes pH 7.5	7.68	437	288	30	1
17	30% PEG 4000 Tris.HCl pH 8.5 0.2 M Li Sulfate	8.96	134	544	70	3
18	20% PEG 8000 Na Cacodylate pH 6.5 0.2 M Mg Acetate	6.62	137	546	72	1
19	30% Isopropanol Tris.HCl pH 8.5 0.2 M NH ₄ Acetate	8.37	218	525	7	0
20	25% PEG 4000 Na Acetate pH 4.6 0.2 M NH ₄ Sulfate	4.95	56	658	37	1
21	30% MPD Na Cacodylate pH 6.5 0.2 M Mg Acetate	6.71	265	468	19	3
22	30% PEG 4000 Tris.HCl pH 8.5 0.2 M Na Acetate	8.96	95	590	65	2
23	30% PEG 400 Na Hepes pH 7.5 0.2 M MgCl ₂	7.28	276	454	25	1
24	20% Isopropanol Na Acetate pH 4.6 0.2 M CaCl ₂	4.64	159	584	11	0
25	1.0 M Na Acetate, Imidazole pH 6.5	7.90	566	166	18	1
26	30% MPD Na Citrate pH 5.6 0.2 M NH ₄ Acetate	6.50	235	509	7	1
27	20% Isopropanol Na Hepes pH 7.5 0.2 M Na Citrate	7.48	280	470	4	2
28	30% PEG 8000 Na Cacodylate pH 6.5 0.2 M Na Acetate	6.89	87	602	65	2
29	1.6 M K ₂ Na Tartrate Na Hepes pH 7.5	7.67	554	180	16	0
30	30% PEG 8000 0.2 M NH ₄ Sulfate	3.84	39	676	38	3
31	30% PEG 4000 0.2 M NH ₄ Sulfate	3.78	147	572	35	0
32	2.0 M NH ₄ Sulfate	5.01	201	517	36	2
33	4.0 M Na Formate	7.68	328	387	36	3
34	2.0 M Na Formate Na Acetate pH 4.6	5.48	274	444	34	2
35	1.6 M K ₂ Na Phosphate Na Hepes pH 7.5	4.52	241	445	15	2
36	8% PEG 8000 Tris.HCl pH 8.5	8.61	371	359	22	5
37	8% PEG 4000 Na Acetate pH 4.6	4.85	192	536	24	1
38	1.4 M Na Citrate Na Hepes pH 7.5	7.95	90	597	62	11
39	2.0 M NH ₄ Sulfate Na Hepes pH 7.5 2% PEG 400	7.67	198	492	63	6
40	20% Isopropanol + 20% PEG 4000 Na Citrate pH 5.6	6.59	157	565	30	0
41	10% Isopropanol + 20% PEG 4000 Na Hepes pH 7.5	7.46	134	560	58	6
42	20% PEG 8000 0.05 M K Phosphate	4.62	89	586	52	3
43	30% PEG 1500	5.36	101	594	54	7
44	0.2 M Mg Formate	6.86	473	253	26	1
45	18% PEG 8000 Na Cacodylate pH 6.5 0.2 M Zn Acetate	5.85	160	572	19	5
46	18% PEG 8000 Na Cacodylate pH 6.5 0.2 M Ca Acetate	6.50	120	570	57	2
47	2.0 M NH ₄ Sulfate Na Acetate pH 4.6	4.64	100	619	30	1
48	2.0 M NH ₄ Phosphate Tris.HCl pH 8.5	4.25	216	461	14	1
Totals			11,102	23,205	1601	99

[†]Buffers when present were at 0.1 M.^apH is experimentally measured pH, performed in duplicate on two different batches of the screen.^bNumber of proteins for which this condition yielded the only crystal lead.

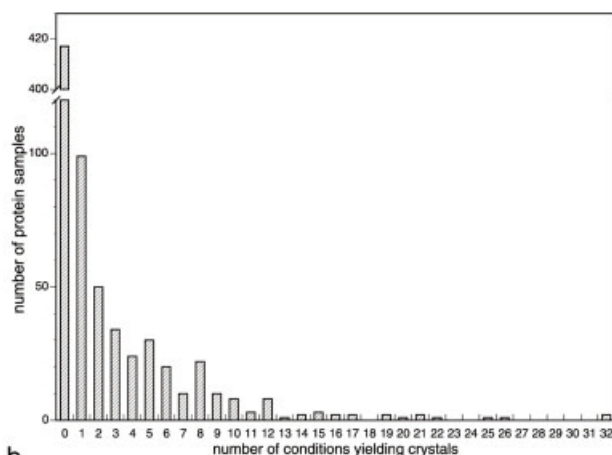
natural host, or may simply reflect the quality of the protein produced by the *E. coli* host. Overall, 338 protein samples (45%) yielded at least 1 crystallization lead. For each protein that crystallized, crystals were observed, on average, in 4.7/48 conditions.

There were large differences in the number of proteins that crystallized in each condition (Table II), ranging from 76 for condition 9 to 4 for condition 27. Surprisingly, for 99 of the 338 proteins crystallized (29.3%) crystals were obtained in only one condition; this is approximately

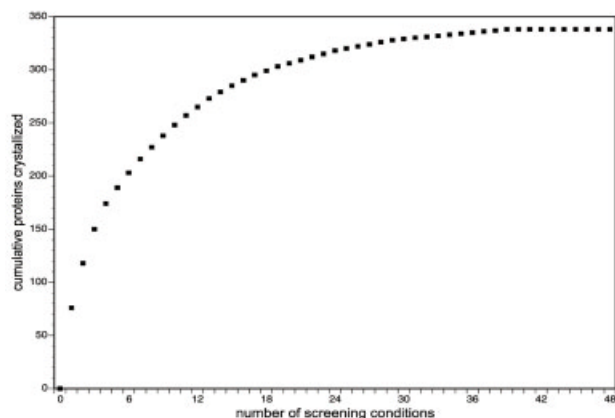
10-fold higher than might be expected if crystallization in different conditions were behaving as independent random variables ($48 \times 0.1 \times 0.9^{47} \sim 3.4\%$).



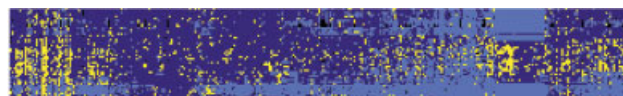
a



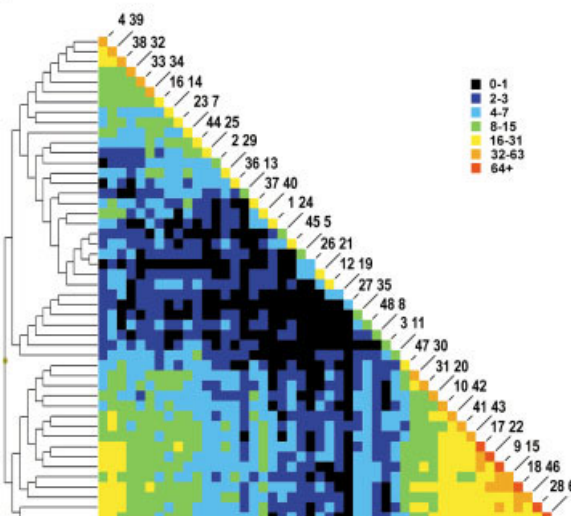
b



c



a



b

Fig. 2. (a) Subset of the overall data, representing the 338 proteins successfully crystallized. Different conditions are arrayed vertically and different proteins horizontally. Yellow squares represent crystals, dark blue squares represent precipitated proteins, and cyan squares represent soluble proteins. Proteins and conditions were sequentially clustered using ClustalX using identity matrices for scoring. (b) Distance matrix of clustered conditions. Condition numbers are noted along the diagonal; off-diagonal elements represent the number of instances in which proteins were found to crystallize in both the condition to the right and the one above it. Note the strong correlation between the PEG conditions (bottom right corner) as well as the citrate/sulphate/formate salts (top left corner). The scale is logarithmic. On the left is a "Cladogram" showing the degree of relatedness of crystal screen conditions as inferred from the degree to which they crystallize the same protein samples. Note that the crystallization conditions cluster primarily on the basis of the chemical nature of the major precipitant and secondarily on the basis of pH.

Efficacy of Different Precipitants

In 17 of the 48 conditions, salt was the major precipitant. These conditions together crystallized a total of 200 of the 338 proteins (59.2%) [Fig. 1(a)]. There were significant differences in the effectiveness of different salts. Sodium citrate, for example, is represented by a single condition in the screen, 39, but was the eighth most productive condi-

Fig. 1. (a) Venn diagram showing the number of proteins for which crystals were obtained in conditions where salt (17 screening conditions, red), PEG (22 conditions, blue), or an organic molecule (9 conditions, green) was the major precipitant. (b) Number of proteins with a given number of successful screening conditions. While some exceptional samples crystallize in up to two-thirds of all conditions, most proteins crystallize in relatively few conditions. (c) Number of crystals contained for a given number of selected screening conditions. Conditions were added one at a time, where the condition added was the one that most increased the number of crystals obtained relative to the previously chosen subset. Note that almost all of the crystals obtained can be obtained from approximately half of the screening conditions and that the last nine conditions could be omitted without affecting the number of samples crystallized.

TABLE III. Minimal Screens[†]

Screen	Conditions included	# proteins crystallized	% crystals vs. full screen	% of total proteins crystallized
Minimal screen 6	6, 10, 18, 38, 39, 43	205	61	27.1
Minimal screen 12	4, 6, 10, 17, 18, 30, 36, 38, 39, 41, 43, 45	268	79	35.5
Minimal screen 24	1, 4, 6, 10, 11, 13, 14, 16, 17, 18, 20, 21, 28, 30, 33, 34, 35, 36, 38, 39, 41, 42, 43, 45	318	94	42.1
JK 1–48	1–48	338	100	44.8

[†]These screens are not only potentially useful in their own right (e.g., in cases where material is limited) but also may potentially serve as the nucleus of a new, more efficient screen.

tion in the screen. Moreover, this condition was by far the most likely condition to yield the only crystal for a given protein. This unusual behavior may be related to citrate's strong metal chelating abilities, something that suggests further investigation.² Ammonium sulphate, long considered an exceptionally good salt for crystallizing proteins, was the major precipitant in four conditions (4, 32, 39, and 47), all of which yield substantial numbers (50, 36, 63, and 30) of crystals. Lithium sulphate (16) and formate salts (33, 34, and 44) were also successful. Citrate, sulphate, and formate salts formed a cluster that affected protein solubility in a similar fashion; they tended to crystallize the same subset of proteins (Fig. 2). The other salts used, namely, phosphate (3, 11, 35, and 48), acetate (7 and 25), and tartrate (2 and 29), were much less successful precipitants; in the case of phosphate, this likely reflected the fact that all of these conditions are acidic. Only 69 proteins crystallized in these 8 conditions.

High-molecular-weight PEGs are widely considered the most successful protein crystallization agents.^{2,9} Of the 338 total proteins, 229 crystallized (67.7%) in the 14 conditions that consist of PEG 4000 or 8000 at concentrations greater than 18%, possibly with some buffer and inert salt (excluding condition 45, which contains zinc), for an average of 54 crystals per condition. The six most productive conditions overall were all in this group. However, there was, in general, a great deal of redundancy among the PEG conditions [Fig. 2(b)]. This region of parameter space appears to be heavily oversampled, a consequence of strong bias toward previously successful experiments built into the original design of the screen. Low concentrations of PEG (36, 37) appeared to be less effective at obtaining crystals but crystallized a different set of proteins than those obtained at higher concentrations. PEG 400 (13, 14, and 23) appeared to affect the solubility of proteins in a manner that more resembled the action of a salt than that of a high-molecular-weight PEG. This may reflect the fact that smaller PEGs probably precipitate proteins more by a solvent competition effect than a volume exclusion effect. Also, zinc in conjunction with PEG yielded different solubility behavior than other salt/PEG combinations, likely reflecting the ability of transition metals to effect strong interactions between proteins by virtue of their ligand chemistry.

Organic precipitants formed a minor part of the screen. There are four MPD conditions (1, 5, 21, and 26) that yielded a modest numbers of crystals (39 proteins crystal-

lized, average of 14 crystals per condition). The 5 conditions in which isopropanol was the primary precipitant (8, 12, 19, 24, and 27) yielded few crystals (average 9 crystals per condition, 39 proteins crystallized) unless combined with 20% PEG (40 and 41). In vapor diffusion experiments, the slow diffusion of water from the drop, where vapor pressure is higher, to the well, where vapor pressure is lower, drives a gradual increase in protein and precipitant concentration in the drop that may eventually lead to the crystallization of the protein. Volatile precipitants, on the other hand, tend to diffuse in the opposite direction, decreasing protein concentration. In the case of isopropanol this happens rapidly, making this precipitant perhaps better suited to batch experiments than vapor diffusion. Overall, the 9 conditions that contain organic precipitants crystallized 66 proteins, 10 of which grew only in these conditions.

Mining the Information to Identify Minimal Screens

The strong interdependence of various conditions of the Jancarik and Kim screen implies that it is, in its present formulation, less than ideal. Several conditions produced few crystals, while other groups of conditions, such as those based on high-molecular-weight PEGs, were too highly correlated. We set out to identify reduced condition sets that minimally compromise the chances of getting at least one crystal. Sequential searches of all combinations of conditions yielded a series of minimal screens (Table III) that comprised sets of conditions optimized for maximal probability of successfully obtaining a crystal [Fig. 1(c)]. A potential drawback of optimizing coverage at the expense of redundancy was that alternate crystal forms with different diffraction qualities might be missed; in those instances where a different crystal form would prove desirable, a second round of screening could then be initiated.

Six conditions—6, 10, 18, 38, 39, and 43 (referred to hereafter as minimal screen 6)—yielded crystals for 205 of the 338 proteins (60.6%) successfully crystallized by the full screen. It is interesting to note the dispersion of these conditions—high-molecular-weight PEG at acid, neutral, and basic pH (10, 18, and 6), low-molecular-weight PEG (43), and two different salts (38 and 39). Augmenting this set with conditions 4, 17, 30, 36, 41, and 45 (minimal screen 12) yielded 268 of the 338 crystals (79.3%), and adding a further 12 conditions—1, 11, 13, 14, 16, 20, 21, 28,

33, 34, 35, and 42 (minimal screen 24)—yielded 318 of the 338 crystals (94.1%). In addition to the conditions defined as minimal screen 24, the omission of conditions 22, 23, 27, 32, and 46 from the screen would have each resulted in 2 fewer proteins being crystallized, while the omission of conditions 8, 9, 12, 15, 25, 26, 37, 44, 47, and 48 would have each resulted in 1 fewer protein being crystallized. Nine conditions—2, 3, 5, 7, 19, 24, 29, 31, and 40—could have been omitted from the screen entirely without losing a single crystal from 755 samples.

Note that had the conditions for the minimal screens been chosen by considering only the total number of crystals produced per condition significantly less productive screens would have resulted. For example, the 6 individually *most productive* conditions crystallized 180 proteins compared to 205 for the optimal 6.

Screening data can also be mined for trends in precipitation. For example, the conditions employing tartrate and acetate salts as the primary precipitant were not only among the poorest crystal producers but also among those least likely to precipitate a protein. Because for the majority of proteins supersaturation was never reached with these precipitants, a substantially higher concentration may be predicted to be more effective.

Rational Strategy for Producing Maximally Productive Screens

Although the Jancarik and Kim screen has proved a useful tool for a generation of crystallographers, it is clear that a more efficient screen of a similar size could be derived from it by substituting some of its less productive conditions with ones chosen that demonstrably complement its more productive conditions. With sufficient data it is a relatively straightforward procedure to eliminate those conditions that contribute little, and iterative cycles of further additions, testing, and elimination should allow the eventual optimization of the screen. Analysis of the data generated may also help suggest suitable candidate conditions for expanding the screen. Conditions that show a strong tendency to uniquely crystallize proteins are likely in regions of parameter space that are under-sampled and could therefore yield more crystals. For example, of the 62 proteins crystallized by condition 38, 11 are uniquely crystallized by this condition, implying that citrate salts have some unique properties whose potential for crystallization is underexploited by the present screen. Similarly, further conditions with transition metal ions (such as condition 45) and intermediate-molecular-weight PEGs (such as in condition 43) might also prove useful additions.

Employing such a strategy, one can experiment with a wide variety of conditions without increasing the amount of work to unmanageable proportions and without sacrificing the proven productivity of a core set of conditions—important considerations given that the only practical manner to obtain sufficient samples and data to implement this strategy is to incorporate it into ongoing structural proteomics efforts. Also, because of the tentative nature of additions, this strategy should encourage the

exploration of chemically diverse conditions that might otherwise not be thought sufficiently “safe” to include in a fixed, generally used screen.

In this study, a crystallization database was created and mined to find the most productive screening conditions, highlighting the value of such efforts within structural proteomics projects. Clearly, not only this sort of information can be extracted from the data, and there is more analysis to perform. For example, whereas this data set comprises the best conditions to identify initial crystals, most of these crystals have not been optimized to form diffraction-quality crystals. It will be interesting to learn whether there are differences in the suitability of different conditions as a starting point for producing large, well-diffracting crystals. The data could also be analyzed for solubility properties rather than crystallization. In this way, one may be able to identify a set of solution conditions that most often yield soluble protein—something potentially useful for NMR experiments, for example. Both solubility data and crystallization data should ultimately be linked to the biophysical properties of the proteins, such as the isoelectric point or amino acid content. Clearly, the crystallization databases resulting from structural proteomics projects can yield important information, and efforts should be expended to ensure that they are routinely collected in a consistent, machine-readable format.

CONCLUSIONS

Initial crystallization conditions for a novel macromolecular sample are obtained by screening the sample against a wide variety of chemical “cocktails,” in general a generic set preselected for their historically proven efficacy in producing crystals. Despite the critical nature of this step, however, no systematic effort has been made to optimize the set of conditions to be used. Here, we used data generated by subjecting a large set of proteins against a commonly used, commercially available screen to obtain a clearer picture of the overall efficacy of the present screening strategies and see if there are obvious ways to improve them. This data leads to several nontrivial conclusions: (1) Among archaeal and bacterial genomes, there appear to be large differences in the degree to which proteins are tractable to crystallization; (2) a small subset of the conditions, even in the relatively small Jancarik and Kim screen, are responsible for a large proportion of crystals obtained overall; (3) as a corollary to this, screening hundreds of conditions, as advocated in some screening protocols, is little more likely to yield a crystal than searching a few tens of well-chosen conditions. The results of this experiment suggest that iteratively adding new conditions, testing against a large set of proteins, and rejecting those conditions that contribute least should allow the fine-tuning of existing screens while still having a useful screen in place at all times. Ultimately this will be of great benefit to structural proteomics efforts as an efficient, optimized screen will give maximal samples for structure solution while minimizing the amount of time and material wasted on unneeded experiments.

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