

Fluorogenic, Subsingle-Turnover Monitoring of Enzymatic Reactions Involving NAD(P)H Provides a Generalized Platform for Directed Ultrahigh-Throughput Evolution of Biocatalysts in Microdroplets

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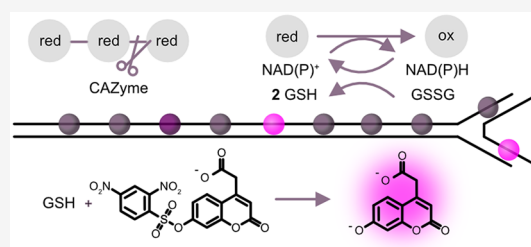


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ABSTRACT: Enzyme engineering and discovery are crucial for a sustainable future bioeconomy. Harvesting new biocatalysts from large libraries through directed evolution or functional metagenomics requires accessible, rapid assays. Ultrahigh-throughput screening formats often require optical readouts, leading to the use of model substrates that may misreport target activity and necessitate bespoke synthesis. This is a particular challenge when screening glycosyl hydrolases, which leverage molecular recognition beyond the target glycosidic bond, so that complex chemical synthesis would have to be deployed to build a fluoro- or chromogenic substrate. In contrast, coupled



assays represent a modular “plug-and-play” system: any enzyme–substrate pairing can be investigated, provided the reaction can produce a common intermediate which links the catalytic reaction to a detection cascade readout. Here, we establish a detection cascade producing a fluorescent readout in response to NAD(P)H via glutathione reductase and a subsequent thiol-mediated uncaging reaction, with a low nanomolar detection limit in plates. Further scaling down to microfluidic droplet screening is possible: the fluorophore is leakage-free and we report 3 orders of magnitude-improved sensitivity compared to absorbance-based systems, with a resolution of 361,000 product molecules per droplet. Our approach enables the use of nonfluorogenic substrates in droplet-based enrichments, with applicability in screening for glycosyl hydrolases and imine reductases (IREDs). To demonstrate the assay’s readiness for combinatorial experiments, one round of directed evolution was performed to select a glycosidase processing a natural substrate, beechwood xylan, with improved kinetic parameters from a pool of $>10^6$ mutagenized sequences.

INTRODUCTION

Enzyme engineering campaigns rely on functional screening for the discovery of starting points and subsequent directed evolution.¹ In light of the increasing demand for biocatalysts in the transition toward a sustainable bioeconomy,² sensitive assays compatible with ultrahigh-throughput screening formats are crucial for overcoming the challenges arising from finding rare functions in the proverbial vastness of sequence space^{3,4} in a time-efficient manner.^{5,6} However, spectroscopic assays routinely used for enzyme screening rely on bespoke model substrates containing aromatic fluorophores or chromophores released as leaving groups upon catalysis,^{4,6–12} but these model substrates rarely emulate the recognition features of many target reactions. Moreover, following directed evolution’s basic law “you get what you screen for”, adaptation of a target enzyme to a model substrate can result in minimal improvements to the desired reaction of a natural substrate.^{13,14} Conversely, by coupling the enzymatic reaction to a reporter reaction by means of a common cosubstrate, such as NAD(P)H, substrates that are neither fluorogenic nor chromogenic themselves^{15,16} may be assayed using fluorescent- or absorbance-based techniques. As any enzymatic reaction dependent on the common cosubstrate can be coupled to such an assay, these are

versatile (“plug-and-play”), with little change in setup required between different classes of enzymes. Beyond their use in enzyme discovery and engineering, these assays may also function as analytical tools for the quantitative detection of small molecules within complex mixtures^{17,18} by virtue of the intrinsic specificity afforded by enzymes. In such a case, the engineerability of reporter enzymes enables the creation of tailored sensors for the desired analyte. In this study, we introduce the novel fluorogenic dinitrophenyl sulfonyl coumarin probe ($S_{N}Ar$ probe For Rapid Assay of NADPH, SAFRAN) in a coupled assay format for quantification of NADH and NADPH-dependent enzymatic activity at ultrahigh throughput. We demonstrate SAFRAN’s ability to assay both NADH-forming and NADPH-consuming reactions and exemplify coupled droplet-based selections for two representa-

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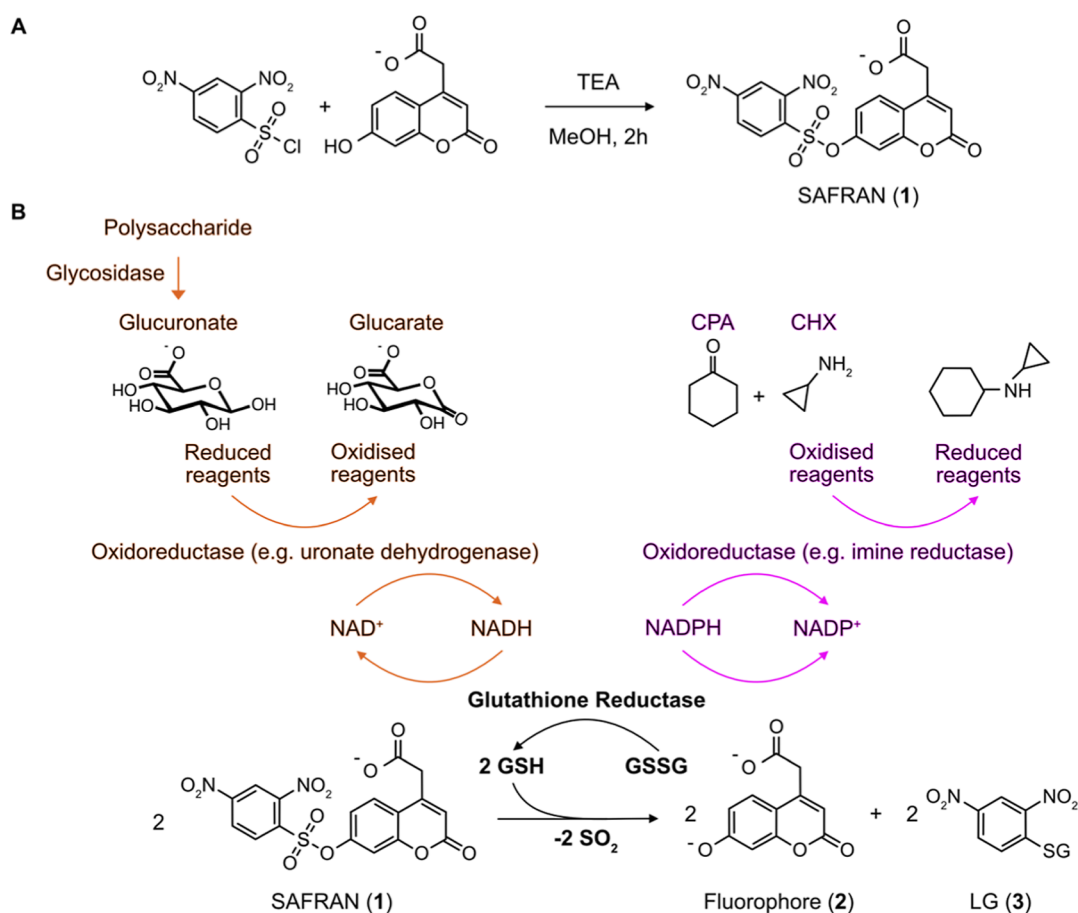


Figure 1. SAFRAN synthesis and coupled assay scheme. (A) Synthesis of SAFRAN 1 from commercially available components; (B) An analyte released by an upstream glycosidase is oxidized (orange) by a specific downstream oxidoreductase, producing or consuming the NAD(P)H that serves as the substrate for glutathione reductase. Reduced glutathione (GSH) spontaneously reacts with SAFRAN 1 forming the coumarin fluorophore product 2 (and the GSH adduct 3). Alternatively, a primary reaction can be detected that modulates NAD⁺/NADPH interconversion (purple). In this scheme, an imine reductase consumes NADPH, which can then be titrated by the downstream detection cascade.

tive enzyme classes: (i) a glycosyl hydrolase linked to a sugar dehydrogenase as a sensor enzyme and (ii) an imine reductase (IRED) that consumes NADPH for the synthesis of chiral amines from ketones.

The reduction of NAD(P)⁺ to NAD(P)H itself can be monitored by an increase in absorbance at 340 nm or vice versa to assay reductive enzymes. This colorimetric change lies at the heart of many small-molecule quantification kits, such as those sold commercially for monosaccharide detection.¹⁹ However, the low molar extinction coefficient of NADH imposes a de facto detection limit of 10 μM in plates using direct absorbance of the cofactor.²⁰ Previous strategies to address these limits have focused on coupling NADH reduction to tetrazolium dye reduction,^{20–22} boosting the measured molar absorption coefficient and improving the signal of the assay 3-fold. Despite these advances, miniaturized ultrahigh-throughput experiments in microfluidic droplets still suffer from a substantial absorbance detection limit of around 10 μM even while using costly dyes.^{5,6,20} Improvements in sensitivity can be achieved using fluorescence assays, as they do not rely on light fully traversing the analyte solution and detect at a wavelength different to the source.²³ While NAD(P)H is itself a fluorophore, its quantum yield is low (2% in aqueous solution, leads to no major sensitivity improvement from using fluorescence compared to absorbance in plate format),^{24,25} and its fluorescence lifetime as well as excitation/emission maxima

are dependent on protein binding,^{26,27} complicating assays in complex mixtures such as cell lysate.^{28,29} Coupling a fluorogenic reaction to NAD(P)H is therefore a compelling alternative, boosting the sensitivity of fluorescent detection while overcoming the limitations of molecular recognition. However, past efforts to do this have been hindered by droplet–droplet leakage of the fluorescent product.³⁰ We achieve this by an enzymatic cascade that detects NAD(P)H and results in the activation of SAFRAN, a profluorescent probe designed to be leakage-free in microfluidic droplets.

The improvement of glycosyl hydrolases is of direct interest for sustainable biocatalysis, and members of the family GH115 are particularly promising in this context as they debranch the hemicellulose xylan, removing internal glucuronic acid residues.³¹ These functional groups are crucial for the recalcitrance of biomass to enzymatic degradation.³² Enzymes to remove these linkages ever more efficiently promise clean ways to utilize the plant cell wall.³³ However, low activity and poor expression have held back industrial applications of enzyme candidates,³⁴ in which cocktail formulations must be finely tuned, while increased enzyme loadings alone are frequently insufficient for optimal saccharification.^{35,36} Therefore, it has been hypothesized that engineering of members of the GH115 family to be more active toward a broader range of substrates will unlock the promise of this family for biocatalytic applications.³⁵ High-throughput assay of the glycosidase

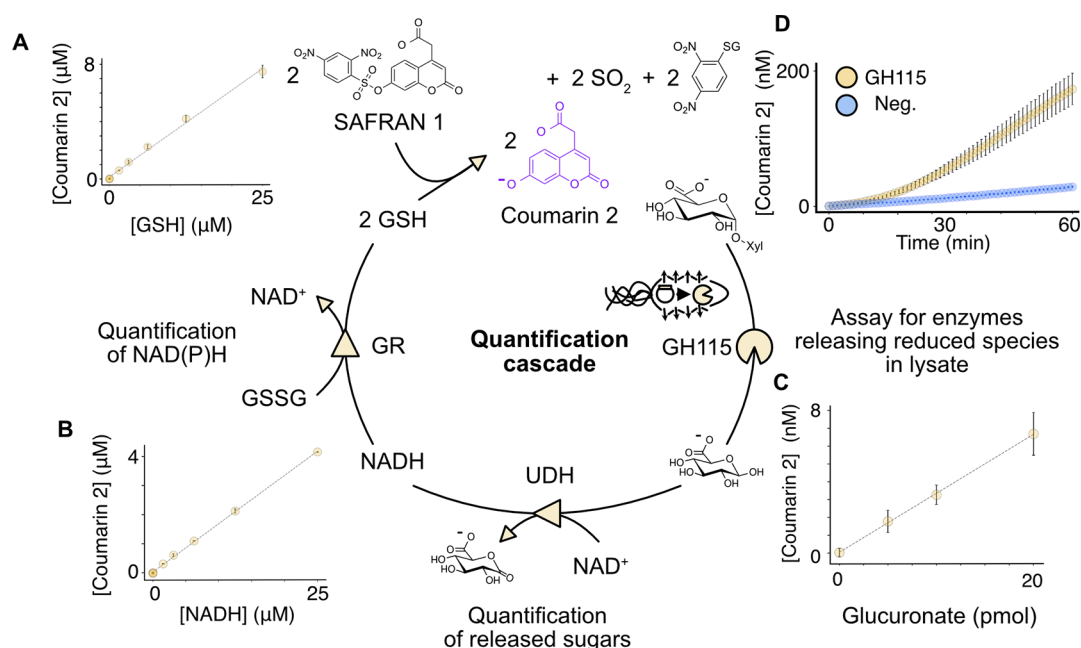


Figure 2. A versatile coupled assay enables detection of trace amounts of monosaccharides in plate format. (A) Unless otherwise stated, all reactions were measured in triplicate at 23 °C, in 200 mM Tris–HCl (pH 7.0), excited at 380 nm and its emission measured at 460 nm. Quantification of fluorescence emission after reaction completion (after 90 min) shows a linear response to reduced glutathione addition. The limit of detection was 50 nM, as defined by a signal significantly above baseline ($p = 0.04$, Welsh's one-sided t -test, $n = 3$ per sample). (B) The S_NAr reaction was coupled to glutathione reductase to consume NADH, and product quantification was achieved based on a linear calibration curve, in this case using fluorescence emission after 90 min of incubation. The limit of detection was 30 nM ($p = 0.001$, Welsh's one-sided t -test, $n = 3$ per sample). (C) The cascade can be further coupled to a monosaccharide dehydrogenase (uronate dehydrogenase) facilitating accurate quantification of glucuronate concentration with a detection limit of 5 pmol per well, or 25 nM ($p = 0.02$, Welsh's one-sided t -test, $n = 3$ per sample). Superior sensitivity was achieved by using 200 mM Tris–HCl (pH 8.0), and over 1 h (the measurement time point), the reaction had reached completion. (D) Time-course of fluorescence emission of GH115-expressing cells with the glucuronoxylan substrate and assay components, compared to cells expressing the control enzyme *SrIRED*, demonstrating that assay can be applied in lysate for quantification of the degradation of complex saccharides or polymers. Xyl = xylan.

reaction has, however, been hampered by the molecular recognition of the substrate routinely extending far beyond the target glycosidic bond itself.^{37,38} Thus, when model substrates lacking specificity-defining features were used to provide an optical readout in directed evolution campaigns after cleavage, the kinetic gains shown by the target enzyme did not translate well to the actual target substrate, as shown in Table S1, creating demand for methods that permit glycosidase screening against natural substrates.^{39–41} We achieve such an assay by linking the release of glucuronic acid residues with NADH formation, and further to SAFRAN, via a specific dehydrogenase acting as a reporter (Figure 1). As an abundance of selective, NAD(P)H-dependent oxidoreductases have been described,⁴² this methodology can be easily expanded to detect a wide range of functional groups and small molecules.

For an application of this new assay in the reverse direction (i.e., for NADH consumption rather than formation), we chose to investigate an enzymatically catalyzed imine reduction reaction, in which NAD(P)H is depleted in a C–N bond forming reaction, for the specific enzyme in question, using the phosphorylated cofactor NADPH instead of NADH. Imine reductases (IREDs) are widely used to access chiral amines, a crucial functionality found abundantly in pharmacologically interesting compounds.⁴³ As most IREDs require engineering before becoming industrially viable and most reactions do not involve strongly fluorogenic compounds, there is a demand for rapid, sensitive assays in the field. As this reaction directly consumes NAD(P)H, SAFRAN can be applied to back-titrate

and quantify NAD(P)H depletion in picodroplets, replacing the less-sensitive tetrazolium dye method.⁴⁴

We benchmark the application of SAFRAN by quantification of NAD(P)H concentrations and, specifically for glycosyl hydrolases, of monosaccharides at ultrahigh-throughput while still matching the detection limit of high-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD), the acknowledged gold-standard column-based method for monosaccharide detection.^{32,45,46} Compared to previously reported coupled assays for NAD(P)H detection in droplets with a low micromolar limit of detection,²⁰ our system provides a 300-fold improvement in sensitivity and maintains the integrity of the product readout over multiple days, enabling extended incubation times.

Having quantified the scope of this assay, we demonstrate its utility in ultrahigh-throughput screening in microfluidic droplets by substantial enrichments in mock library selections (from defined plasmid mixtures) for both example reactions, glycosidases and IREDs, in forward and reverse assay directions. Finally, we apply the SAFRAN cascade in directed evolution, starting from a library of mutant GH115 glycosidases, and screen >10⁶ members for glucuronoxylan-debranching activity. We identify and recover an abundance of mutants with improved lysate activity and, after purification of five of the mutants, identify a sextuple mutant with a 2-fold increase in k_{cat} . Based on these findings, we propose that the SAFRAN-coupled assay represents a widely applicable method

for enzyme discovery, directed evolution, and small-molecule quantification in picodroplets.

RESULTS AND DISCUSSION

NAD(P)H-Based Detection Cascade Based on Thiol-Dependent Uncaging of SAFRAN. To couple NAD(P)H production with turn-on fluorescence, we designed a cascade linking the emergence of NAD(P)H to the formation of free thiols via enzymatic glutathione reduction. The free thiols subsequently uncage a synthesized fluorogenic probe (SAFRAN) in a rapid and irreversible S_NAr reaction with a dinitrophenol group that was quenching coumarin fluorescence.^{47–50} Synthesis of SAFRAN **1** was readily achieved by condensation of 7-hydroxycoumarinyl-4-acetic acid **2** with 2,4-dinitrobenzenesulfonyl chloride in one step from commercially available compounds at 76% yield without the need for purification by chromatography (Figure 1A). High sensitivity is ensured with a favorable reaction stoichiometry: two equivalents of reduced glutathione are produced for each equivalent of NAD(P)H consumed. Any NAD(P)H-producing enzyme could in principle be added to this cascade to elicit a fluorogenic response (Figure 1B). The cascade reagents are cheap and generally accessible; NAD(P)H is reoxidized by the commercially available glutathione reductase, so the expensive NAD(P)⁺ cofactor only needs to be present in catalytic quantities.

The design of SAFRAN was focused around a strongly fluorescent 7-hydroxycoumarin core—as coumarin derivatization is robust and well established, the probe could be readily functionalized.^{51,52} Leakage is a well-documented challenge in microfluidic assay development, preventing the use of the typically large hydrophobic and aromatic probes for all but the fastest reactions.^{53–57} This poses a hitherto unsolved challenge for fluorogenic NAD(P)H quantification as resorufin, the sole published fluorogenic probe for NAD(P)H detection,^{54,58} is hydrophobic and consequently only compatible with very short reaction times in microfluidic droplets.⁵⁹ To minimize the leakage of SAFRAN and its fluorophore product **2**, a carboxylic acid group was incorporated to impart a negative charge on the coumarin scaffold at physiological pH, enabling application in droplet microfluidics with minimal leakage.

As the fluorescence of the umbelliferone structure is dependent on its protonation state (determined by the pH of the reaction medium⁶⁰), we obtained fluorescence excitation and emission spectra of the 7-hydroxycoumarin fluorophore under different pH conditions (Figure S1). Under physiological to slightly basic conditions, the highest signal intensity was achieved with an excitation wavelength of 380 nm coupled with an emission wavelength of 460 nm.

Detecting Picomoles of Monosaccharides in High Throughput. To determine the sensitivity of SAFRAN **1** to reduced glutathione (GSH), GSH was added across a concentration range (25 nM to 25 μ M), eliciting a linear response over three orders of magnitude after 90 min of incubation (Figures 2A and S2). Formation of phenylated glutathione product **3** and release of coumarin fluorophore **2** were confirmed via mass spectroscopy (Figure S3). Subsequently, we demonstrated that glutathione reductase couples the presence of NADH to the S_NAr reaction and thus fluorogenic signal (Figure 2B), with a detection limit of 30 nM NADH (corresponding to an amount of 3 pmol NADH, $p = 0.001$, Figures 2B and S4) in plates. Taking advantage of the bispecificity of glutathione reductase for both nicotinamide

cofactors NADH and NADPH,⁶¹ we furthermore show the applicability of the SAFRAN cascade for quantification of NADPH in plate format (Figure S5).

To demonstrate the use of the SAFRAN assay as an analytical tool with high sensitivity, we set out to show its potential for improving monosaccharide detection limits with high throughput. Monosaccharide liberation is often associated with the breakdown of complex sugars. However, the most sensitive monosaccharide quantification currently, HPAEC-PAD, requires minutes per single assay, very clean samples,⁶² and large amounts of costly and sometimes unstable standards.⁶³ In contrast, high-throughput plate-based detection of monosaccharides has sensitivity in the micromolar range at best and can suffer from low selectivity for the monosaccharide of interest.⁶⁴ Compared to HPLC-based systems or direct chemical detection, dehydrogenase enzymes are engineerable⁶⁵ and a suite of these enzymes are already available as coupling agents,⁴² so that tailored sensors for any analyte or application can be created.

Using the coupled assay in plate format and selecting uronate dehydrogenase as a representative coupling enzyme, we were able to detect glucuronic acid as a linear signal with a detection limit of 25 nM ($p = 0.02$, Figures 2C and S6), representing only 5 picomoles of analyte per well. This is a >100-fold improvement on the best commercially available plate-based glucuronic acid quantification method¹⁹ and comparable to the 12.5 picomole/sample limit of quantification of HPAEC-PAD.⁶⁶ In addition to fluorescence end-point measurements, monosaccharides could also be quantified within minutes of reaction initiation using a maximum rate method, enabling shorter analysis times (Figure S6). This indicates that the method is a rapid, accessible, and derivatization-free alternative to column-based methods for trace monosaccharide quantification, promising to reduce the equipment and reagent costs involved in sensitive monosaccharide detection.

Small-molecule quantification is a widely used analytical method for quantifying polymer degradation.⁵⁸ Upon confirming that the SAFRAN assay was well-suited for quantifying small molecules, we set out to test the capacity of the cascade to quantify the activity of glycosidases that debranch hemicellulose polysaccharides using *Escherichia coli* as the enzyme expression host. In a cellular context, detection cascades risk suffering interference, both optically and chemically. To test the theoretical limits of detection of the cascade in *E. coli* lysate that may be in place due to optical interference, we diluted coumarin **2** in a working concentration of *E. coli* lysate and determined the limits of detection to be 8 nM coumarin **2** ($p = 0.0001$, Figure S7).

We then set out to evaluate whether chemical interference in the cascade would preclude screening for catalysis in an *E. coli* host. If background activities in the screening host are too high, excessive background signal can make quantification of the target activity impossible. This would reduce the capacity of the cascade to detect enzyme activity in cell lysates, thereby limiting its use for high-throughput enzyme screening. To test the signal-to-noise ratio of this cascade, we incubated *E. coli* preinduced for expression of a glucuronoxylan-debranching enzyme (*AxyAgu115A*) with lysis agent, cascade components, and the glucuronoxylan substrate over an hour-long window. This revealed that following a brief lag-phase, the fluorescence intensity increased up to 4-fold above the negative lysate control within an hour, following glucuronic acid release from

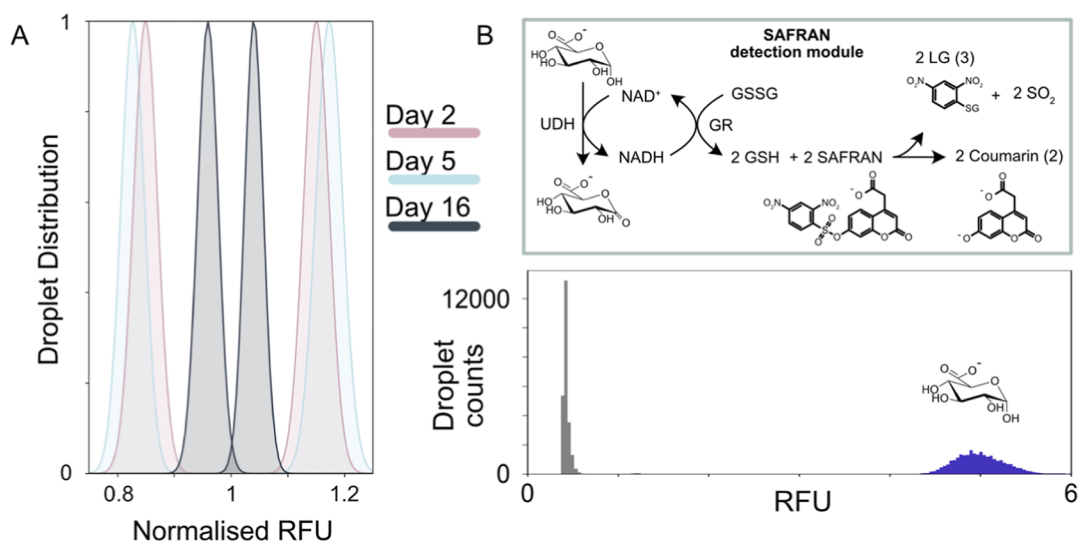


Figure 3. Droplet application of the coupled assay yields a stable signal that emerges rapidly. (A) Co-incubation of droplets containing 25 μM or 50 μM coumarin 2 for up to 16 days maintained two distinct fluorescent populations when measured on-chip. The data shown are the Gaussian fits to the mixed populations, when normalized by RFU and abundance for clarity of presentation. (B) Droplets with cascade reagents and buffer form two distinct populations on a FACS sorter when mixed, incubated for 40 min, and measured together. Despite the high analyte concentration, no droplet-to-droplet exchange of the product (or intermediates) of the reaction cascade that would jeopardize the distinction between positives and negatives is observed.

the polymer (Figure 2D). The ability to use our coupled assay for screening of enzymes in *E. coli* lysate integrates it into a preferred format for plate and droplet-based enzyme engineering and metagenomic screening campaigns.⁶⁷

To demonstrate the broad applicability of the SAFRAN cascade, we also explored its ability to report on a NADPH-dependent reduction reaction, thus highlighting both the tolerance of glutathione reductase for phosphorylated and nonphosphorylated nicotinamide and the capability to use this assay for back-titrations. To this end, the condensation reaction between cyclopropylamine (CPA) and cyclohexanone (CHX) via reductive amination, catalyzed by an imine reductase from *Streptosporangium roseum* (SrIRED) (Figure 1B), was examined. After reductive amination using lysate containing SrIRED, the cascade was added to back-titrate IRED activity, showing a clear time-dependent signal (Figure S5).

Detection of Oxidoreductases in Picodroplets Using SAFRAN. Much larger numbers of protein variants can be screened in an attractive ultrahigh-throughput format where water-in-oil emulsion droplets act as reaction compartments made and handled in microfluidic devices.⁶ However, this format has specific limitations that must be overcome, e.g., the leakage of aromatic, hydrophobic fluorescent probes between droplets, likely due to enhanced oil–probe hydrophobic interactions.⁵⁶ This leakage results in a loss of signal over time, shortening maximum droplet incubation times and thus precluding the detection of enzymes that are poorly expressed or have low activity. Therefore, SAFRAN was designed to carry a net negative charge at physiological pH in the form of a carboxylic acid moiety. Indeed, fluorescence microscopy imaging of a mixed population of droplets, containing either buffer only or 100 μM coumarin 2, showed no equilibration between droplets after 24 h (Figure S8). Further, incubation of a mixed population of droplets containing 25 and 50 μM fluorophore revealed no detectable leakage for up to 5 days (Figure 3A) when measured using fluorescence-activated

droplet sorting (FADS). Measuring the same droplets again at day 16 showed that the fluorescence of the two populations had begun to converge (by 73% of their original fluorescence). However, the individual populations could still be distinguished with only 3% overlap. This indicates a very slow rate of leakage, perhaps inevitable due to surfactant-mediated transfer between droplets.^{68,69} These measurements indicate that incubations of up to at least 5 days should be feasible without facing challenges with leakage. This stands in contrast to the NAD(P)H probe resorufin—fluorescence microscopy in a previous study revealed near complete equilibration of droplets with and without fluorophore in only 6 h,⁵⁹ suggesting SAFRAN achieves at least 64-fold better signal retention over this critical window used in picodroplet screening for medium-to-slow reactions (Table S2).^{7,9,70}

We then set out to test if oxidoreductase activity could be detected in droplets using SAFRAN by again creating a mixed two populations of droplets: one containing the SAFRAN cascade reagents, the oxidoreductase uronate dehydrogenase, and glucuronic acid; the other identical but without glucuronic acid. Fluorescence was present at the limit of detection 10 min after droplet formation (Figure S9) but was distinct from background within 40 min (Figure 3B). Furthermore, these populations remained distinct after incubation for an overnight period (Figure S9). These results demonstrate that the SAFRAN cascade can be used to detect the purified oxidoreductase uronate dehydrogenase in droplets, forming the basis for using these enzymes as coupling agents to detect upstream chemical reactions, such as the release of monosaccharides from polysaccharides.

Sensitivity is a major limitation of several droplet-based detection modes as, for example, absorbance-based detection (~ 10 μM),^{20,71} which currently is the only method available for screening dehydrogenase activity using nonmodel substrates at ultrahigh throughput that does not suffer from leakage.^{59,70} Fluorescent assays can be up to 3 orders of magnitude more sensitive, as shown by Colin et al. for a

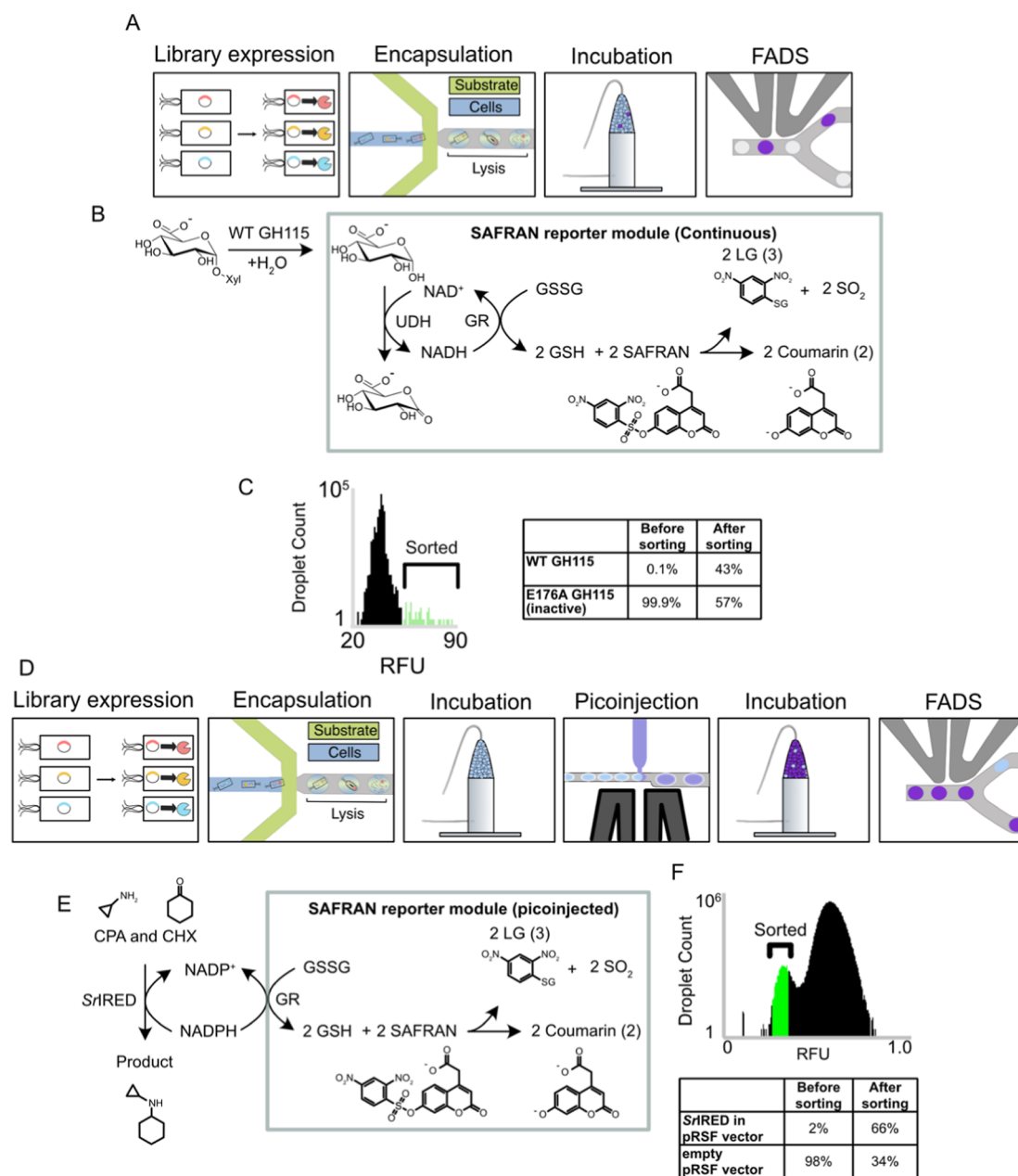


Figure 4. A sensitive droplet assay detects glycosidase and imine reductase activity from single cells on natural substrates. (A) Workflow used for testing the enrichment of glycosidases in droplets. (B) Schematic of the target droplet reaction in this assay. (C) (Left) Width-gated histogram describing a sample of droplet RFUs in the unsorted library. The top 0.02% of droplets were selected for sorting. (Right) Plate activity of sorted clones picked from the recovered library relative to the wild type. 39 of the 90 clones assayed were found to be confidently positive. (D) Workflow used for testing the enrichment of imine reductases in droplets. (E) Schematic of the target reaction in droplets. The imine reductase (IRED) first consumes NADPH until the droplets are picoinjected with the SAFRAN reporter module, which quenches the reaction and converts the unreacted NADPH to a fluorescent signal. (F) Droplet trace and enrichment statistics of the IRED. Post-sorting library composition was determined using Sanger sequencing from 12 recovered clones.

fluorescein-based phosphotriesterase assay with a detection limit of 2.5 nM.⁴ This sensitivity allows detection of enzymatic activities with less than one turnover per enzyme molecule, thus enabling ultrahigh-throughput screening even when expression or intrinsic activity is low. To test the limit of detection of the SAFRAN system using FADS, we sequentially measured the signal from droplets containing different concentrations of coumarin product **2** and found a linear dependence of fluorescence and product concentration with a clear signal detectable down to 30 nM (corresponding to 361,000 molecules per 20 pL droplet, or 180,500 turnovers of

glutathione reductase, Figure S10). The lowest quantified droplet populations tested yielding distinct populations were 0 and 30 nM (Figure S10). A single cell compartmentalized in a droplet typically produces $\sim 10^6$ molecules of enzyme,²⁰ providing a resolution that is able to detect fewer product molecules than the number of expressed enzyme molecules (Figure S10). The assumptions made for this calculation are provided in Table S3, including estimates of the background deriving from the pool of endogenous *E. coli* reduced glutathione. This glutathione pool varies over the course of *E. coli* growth,⁷² and so, it is important to ensure that all

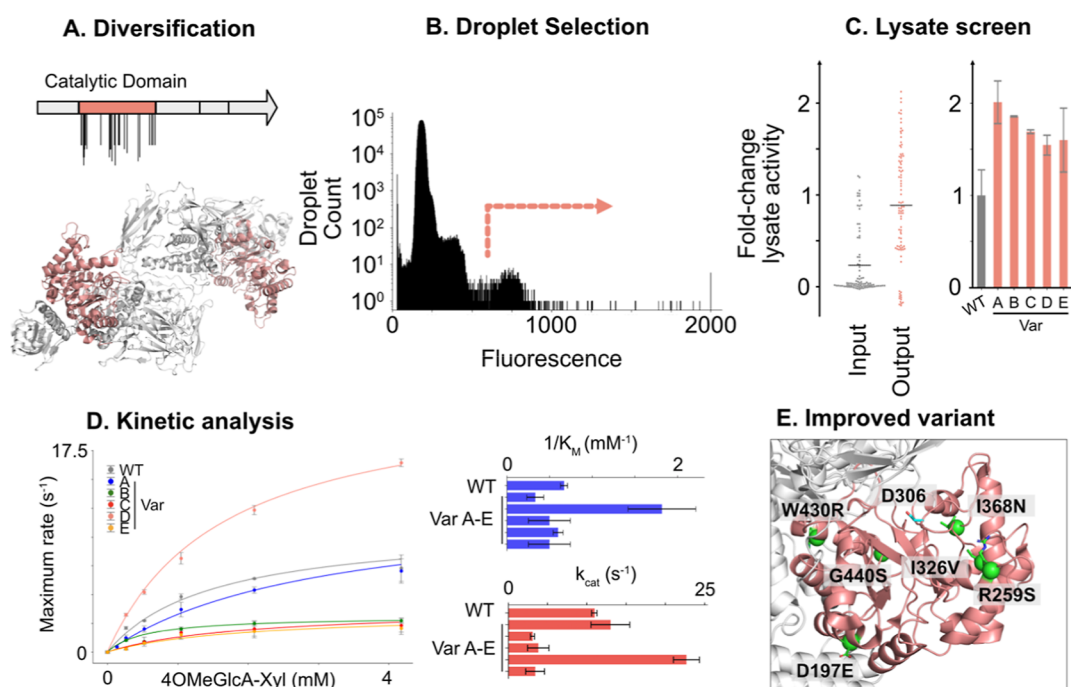


Figure 5. Directed evolution of GH115. (A) Schematic showing the dimeric structure of *AxyAgu115A* (PDB 6NPS), highlighting the catalytic domain. (B) Droplet trace from the sort of 3.6 million droplet events. (C) (Left) Maximal rate of release of 4-OMeGlcA from BWX from the sorted lysate from either the unsorted or sorted library in microtiter plates. Each point has been normalized between WT (=1) and inactive E176A (=0). (Right) Maximal rate of release of 4-OMeGlcA from BWX of the selected improving members. Values represent duplicate growth and measurement, except for wild type, which has six replicate growths and measurements. (D) Kinetic analysis of mutants identified with improvement in lysate. Measurements are the average of three replicates. (E) Schematic showing the position of mutations in variant D (green) relative to a putative catalytic residue (D306, cyan). Structure derives from PDB 6NPS.

growth phases are synchronized to minimize background phenotypic variation.

Screening Plasmid Libraries of Hydrolases and Oxidoreductases against Nonmodel Substrates Using FADS. Following confirmation that the SAFRAN system was amenable to extended droplet incubation, we investigated the use of the system for screening enzyme libraries at ultrahigh throughput in an *E. coli* host. First, we tested the capacity of the system to identify glycosidases releasing monosaccharides, which served as the oxidizable substrate that was coupled to the SAFRAN cascade using a monosaccharide dehydrogenase. The two-step microfluidic workflow tested here is outlined in Figure 4A. This workflow requires expression of the cytoplasmic enzyme library in bulk media followed by coencapsulation of cells with the substrate, coupled assay, and lysis reagents in a flow-focusing device. The droplets were then subjected to an incubation over a suitable time frame to reach the limit of detection (at least 271,000 enzyme turnovers), followed by sorting using FADS. To test this workflow, a defined library consisting of a 1000:1 abundance of inactive GH115 E176A (the relative activities of the WT and E176A are shown in Figure S11) against a minority of wild-type enzyme was screened for activity against a natural substrate, beechwood xylan (420 μ M 4OMe-GlcA motif), as shown in Figure 4B. This library was encapsulated using a flow-focusing device and droplets were stored off-chip for 60 min, after which the most fluorescent 0.02% of droplets were sorted at 800 Hz using FADS (Figures 4C and S12), yielding an enrichment of 765-fold, calculated according to the method of Baret et al.⁷³ (Figure S13, Note S1).

To demonstrate quantification of the backward reaction in droplets, where NADPH is consumed, the SAFRAN assay was

used to enrich for cells expressing the bacterial imine reductase SrIRED catalyzing the condensation of CPA and CHX using the workflow outlined in Figure 4D. Individual bacteria containing either empty plasmid or plasmid containing the IRED gene were allowed to express the IRED, where present, in bulk media before mixing in a 98:2 ratio and coencapsulation together with substrates and lysis reagents using a flow-focusing chip. After subsequent 1 h off-chip incubation time, the droplets were picoinjected with the SAFRAN cascade components, which reported on the amount of unreacted NADPH remaining in each droplet (Figure 4E). Sorting with FADS for the low fluorescence population enabled an 98-fold enrichment of IREDs (Figure 4F) according to the method of Baret et al.⁷³ (Supplementary Note 2).

The successful enrichments of enzymes from the hydrolase and the oxidoreductase reaction classes, using either the cofactor NADH or NADPH, measuring either cofactor production or consumption, establish SAFRAN as a useful method for assaying any reaction that can be coupled to NAD(P)H consumption or production with extremely high sensitivity and at ultrahigh throughput.

Ultrahigh-Throughput Directed Evolution of GH115 against the Feedstock Beechwood Xylan. The SAFRAN assay delivers a promising system for performing directed enzyme evolution against the substrate of choice. The laboratory evolution of GH115 for higher activity against its substrate beechwood xylan promises direct applications in bioreactors for xylan utilization. In particular, improving kinetic parameters of the enzyme permits lower enzyme loadings and more efficient operation in complex mixtures of the substrate. The enzyme used as a starting point, *AxyAgu115A*, is known to

Table 1. Summary of Purified Enzyme Parameters Derived from Five Selected Variants

mutations	relative lysate activity	$k_{\text{cat}}/\text{s}^{-1}$ (fold change)	K_{M}/mM (fold change)	$k_{\text{cat}}/K_{\text{M}}/\text{M}^{-1} \text{ s}^{-1}$ (fold change)
WT	1.0 ± 0.30	10.97 ± 0.34	1.49 ± 0.09	7360 ± 500 (1.00)
A L322Q_Y437F	2.0 ± 0.33	13.02 ± 2.48 (1.19)	2.98 ± 0.87 (2.00)	4400 ± 1520 (0.60)
B L426P	1.9 ± 0.01	3.11 ± 0.27 (0.28)	0.55 ± 0.12 (0.37)	5650 ± 1330 (0.77)
C E253D	1.7 ± 0.03	3.84 ± 1.34 (0.35)	2.00 ± 0.98 (1.34)	1920 ± 1160 (0.26)
D D197E R259S I326V I368N W430R G440S	1.7 ± 0.15	22.68 ± 1.65 (2.07)	1.66 ± 0.16 (1.11)	13,700 ± 1650 (1.86)
E F358L	1.5 ± 0.49	3.43 ± 1.20 (0.31)	2.00 ± 0.98 (1.34)	1715 ± 1030 (0.23)

be a more tolerant enzyme to xylan substitutions than some other GH115 enzymes⁷⁴ and possesses some alkaline tolerance.³³ Alkaline conditions are preferable in xylan processing, as the solubility of the polymer is greater, but making enzymes compatible with these conditions remains a challenge.³³ Therefore, *AxyAgu115A* serves as a promising starting point for the directed evolution of a more efficient GH115 enzyme.

For several reasons, the GH115 family is a potentially challenging target for directed evolution: the enzymes are dimers of around 1000 amino acids per monomer and consist of 4–5 domains, with the second domain “B” acting as the catalytic domain.⁷⁴ Although there is evidence that the enzyme acts via a Koshland inverting mechanism,⁷⁵ the catalytic acid and base have not been unambiguously identified. This is partially because the active sites are, unusual for a glycosidase, composed of flexible loops, hindering crystallographic identification of the catalytic residues. The other domains are of unknown function but are likely catalytically relevant as the active site alone domain is not sufficient for catalysis.⁷⁶

We hypothesized that mutagenesis focused on domain B could target catalytic improvements, while the other four domains were left constant to retain protein stability and any other important yet cryptic roles. Whole-domain mutagenesis was achieved using error-prone PCR (Figure 5A). The substrate concentration in droplets was 420 μM of the (4-OMe)GlcA epitope, almost an order of magnitude below the K_{M} of the enzyme (1.49 mM), thus enacting selection pressure for variants that are able to strongly bind the substrate, targeting improved performance under bioreactor conditions where the epitope comprises only a small fraction of the total target plant cell wall loading. Following off-chip incubation of the library members and xylan for an hour, 3.6 million droplets were analyzed, and the highest fluorescent droplets were sorted (Figure 5B). 39 wells from the output were identified as having higher than wild-type activity in cell lysate (43%, Figure 5C), displaying up to 2-fold improvements in the maximal release rate of (4-OMe)GlcA. Five members A–E were selected from the secondary screen for further characterization.

These mutants were found to harbor up to six mutations (Table 1) and were isolated for further characterization. L426P showed an almost 3-fold reduction in the K_{M} of the enzyme, while the mutant D197E R259S I326 V I368N W430R G440S showed a 2.1-fold increase in k_{cat} and a nonsignificantly changed K_{M} , resulting in 1.9-fold increase in $k_{\text{cat}}/K_{\text{M}}$ (Table 1, Figure 5D). This variant contains mutations scattered around the catalytic domain, with position R259 notably interacting with the disordered loop, which is shown to be critical for catalysis (Figure 5E). By circumventing the dependence on modified substrates, this single round of directed evolution via a SAFRAN-coupled assay achieves among the highest reported improvements in kinetic parameters of glycosidases against their unmodified substrates (Table S1).

DISCUSSION

Our ultrahigh-throughput fluorogenic NAD(P)H assay enables large-scale screening of enzyme libraries against an unmodified substrate of interest, which we have applied here for enzyme engineering by directed evolution. We achieved this by designing a caged coumarin, SAFRAN, that is uncaged following NAD(P)H-dependent reduction of oxidized glutathione and subsequent reaction with the free thiols of reduced glutathione. The coumarin product carries a carboxyl group, preventing leakage between the droplets. We demonstrate that screening in an *E. coli* expression host is compatible with this cascade, with the assay signal overcoming any interference derived from endogenous *E. coli* reduced glutathione and NAD(P)H reserves.

We use the cascade to meet the challenge of engineering glycosyl hydrolases, which alongside medical applications⁷⁷ are an important class of enzymes for their ability to convert biomass into valuable small molecules with high specificity and selectivity.⁷⁸ The enzyme *AxyAgu115A* has recently been used in a cascade reaction to convert hardwood glucuronoxylan, an underused fraction from biorefineries, to the dicarboxylic acid 4-*O*-methyl *D*-glucaric acid.⁷⁹ Glucuronic acid and its derivatives are valuable chemicals for synthetic purposes and demand currently for these compounds far outstrips the current capacities of chemical synthesis and extraction from natural sources.⁸⁰ Glucuronic acid is also being used to phase out less sustainable chemicals in manufacturing; dicarboxylic acids are valuable building blocks for the synthesis of pharmaceuticals and bioplastics, but demand is currently mostly met using petrochemicals.⁸¹ The approach described in this work enabled us to select *AxyAgu115A* variants for their activity against their natural substrate, the feedstock glucuronoxylan, and not against an activated model substrate, thus ensuring that improvements made during the directed evolution translate to industrially relevant catalytic improvements against the polymer of interest.

The application of this assay to insoluble substrates in microfluidic droplets—common targets of interest for industrial applications⁸²—remains to be demonstrated. Unlike the soluble substrates tested in this work, insoluble particles pose challenges of (i) sedimentation, (ii) optical interference with fluorophores, and (iii) destabilization of the oil–aqueous interface. One solution already in use to overcome these challenges are microscale hydrogel beads.⁸³ Hydrogels immobilize particles such as insoluble substrates and even whole cells⁸⁴ and are compatible with fluorescence detection methods.⁸⁵ The immobilization of the substrate in the hydrogel stabilizes the surrounding droplet–oil interface. Hydrogel formats therefore pose an attractive avenue to expand the SAFRAN-coupled assay to screen insoluble substrates.

The modest rate accelerations achieved by directed evolution must be evaluated in the context of the engineering

challenges in the GH115 family. Family members are large dimers with each monomer containing up to 1000 amino acids. Furthermore, beyond dimerization, functions of the domains other than the TIM-barrel catalytic domain have yet to be elucidated. Despite the family seemingly operating an inverting Koshland mechanism,⁷⁵ the active site residues have also not been decisively identified and are likely present on the long and flexible loops that cover the active site and have been implicated in catalysis and substrate scope.⁷⁴ Using the available information on the GH115 family, we focused mutagenesis on the active site domain and found that the most improving variant had mutations not located in the active site but in the second shell residues and in more distant positions throughout the catalytic domain. The position R259, mutated to serine in the improved mutant, is a second-shell residue known to form a salt bridge with the long active site loop and thought to influence the tolerance of the enzyme to substitutions on the xylan backbone.⁷⁴ W430 (mutated to arginine) is found at the domain interface of the catalytic domain and two other domains within the protein, potentially influencing the long-range protein dynamics.

While we demonstrate the cascade for engineering *AxyAgu115A*, the development of the SAFRAN cascade was undertaken with the primary goal of making dehydrogenase-coupled assays amenable to droplet screening. Based on the assay's modularity, we propose that other dehydrogenase-coupled assays can now be analyzed using FADS (or, after a second emulsification step, by standard flow cytometric sorting in double emulsions^{86,87}), taking advantage of the wealth of literature that reports on coupling reactions to NAD(P)H production or depletion. These include xylosidases, glucosidases, arabinosidases,⁷⁰ small-molecule/natural product methyltransferases,⁸⁸ nucleotide methyltransferases,⁸⁹ ATPases and other enzymes that produce ADP^{90,91} including kinases,⁹² systems that produce urea or pyruvate,⁹³ amino acid oxidative deaminases,⁹⁴ phosphite dehydrogenases,⁹⁵ and alkane hydroxylases.⁹⁶ Additionally, engineering of redox enzymes that are not in a coupled setup, such dehydrogenases for biocatalytic applications,^{97,98} including conversion of monosaccharides to hydrogen gas,⁹⁹ will directly benefit from this sensitive screening assay. As a proof-of-principle, we demonstrate how the SAFRAN assay is suitable for screening libraries of imine reductases, a pharmaceutically valuable synthetic family of enzymes.

Based on these degrees of freedom, SAFRAN is a uniquely modular detection system, where the same assay reagent can be deployed in various contexts, the elements of which can be modularly exchanged: (i) On the one hand, a wide range of primary substrates can be detected, as long as they lead to an intermediate that is processed by the coupled reaction. For example, reactions of a wide range of natural glycosides can be detected, as long as monosaccharides that feed into a coupling reaction. Therefore, the range of options for substrate detection is determined by the availability of specific upstream coupling enzymes that can be modularly exchanged. (ii) On the other hand, the reaction type can be chosen by employing any of the above-mentioned enzymes that consume or generate NAD(P)H, here by exchanging the downstream coupling enzymes that process the intermediate substrate in redox reaction.

CONCLUSIONS

In contrast to alternative NADPH detection strategies, the improved sensitivity of our approach²⁰ and reduced leakage between droplets³⁰ allows the detection of low-activity enzymes by extended droplet incubation times. This is crucial for protein engineering and discovery of other biocatalysts,^{100–102} as the recruitment of proteins for the conversion of our current economic model to a sustainable bioeconomy depends on three sources with typically weak activities. These are (i) promiscuous side activities of existing enzymes,^{103,104} (ii) enzymes from metagenomic sources,¹⁰⁵ identified by functional screening or by bioinformatics, and (iii) computationally designed enzymes.^{100–102} Without a sensitive assay, these useful, but imperfect, starting points cannot be improved by directed evolution, and compatibility with an ultrahigh-throughput method is crucial for success. Droplet microfluidics is arguably one of the most powerful UHT formats, allowing screening of $\sim 10^7$ -membered libraries in a day, and SAFRAN makes sensitive fluorogenic screening of a broad range of reactions possible in this format.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/jacs.4c11804>.

Supporting Information and methods (synthesis, protein expression and purification, microfluidic procedures for enrichment and directed evolution, library synthesis, kinetic analysis; supplementary figures (assay calibration curves and sensitivity; supplementary tables (survey of glycosidase evolution outcomes, survey of fluorescence-activated droplet sorting studies, analysis of assay sensitivity); fluorescence spectra of the fluorophore, NMR spectra, mass spectrometric product analysis; fluorescence microscopy images of droplets; nucleotide sequences of enzymes; and supplementary note on enrichment analysis (PDF).

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Notes

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