Whole-cell screening of oxidative enzymes using genetically encoded sensors†

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Biocatalysis is increasingly used for synthetic purposes in the chemical and especially the pharmaceutical industry. Enzyme discovery and optimization which is frequently needed to improve biocatalytic performance rely on high-throughput methods for activity determination. These methods should ideally be generic and applicable to entire enzyme families. Hydrogen peroxide (H₂O₂) is a product of several biocatalytic oxidations and its formation can serve as a proxy for oxidative activity. We designed a genetically encoded sensor for activity measurement of oxidative biocatalysts via the amount of intracellularly-formed H₂O₂. A key component of the sensor is an H₂O₂-sensitive transcriptional regulator, OxyR, which is used to control the expression levels of fluorescent proteins. We employed the OxyR sensor to monitor the oxidation of glycerol to glyceraldehyde and of toluene to o-cresol catalysed by recombinant E. coli expressing an alcohol oxidase and a P450 monooxygenase, respectively. In case of the P450 BM3-catalysed reaction, we additionally monitored o-cresol formation via a second genetically encoded sensor based on the phenol-sensitive transcriptional activator, DmpR, and an orthogonal fluorescent reporter protein. Single round screens of mutant libraries by flow cytometry or by visual inspection of colonies on agar plates yielded significantly improved oxidase and oxygenase variants thus exemplifying the suitability of the sensor system to accurately assess whole-cell oxidations in a high-throughput manner.

Introduction

Several enzyme class 1 (EC 1) members enable highly selective oxidations and are increasingly perceived as advanced catalysts for chemical production. The catalytic properties of oxidative enzymes are undoubtedly unique, yet the performance of the enzyme is often insufficient for preparative synthesis. However, enzyme features such as operational stability, reaction rates, and substrate specificity can be tailored in rounds of random or (semi-) rational diversification of amino acid sequences followed by activity screening of the generated protein sequence space. As a rule of thumb, in protein engineering campaigns the likelihood of discovering an improved variant is proportional to the total number of variants tested. Thus, detection methods with both high discriminatory power and compatibility with high-throughput screening workflows are essential for the successful discovery and engineering of industrial biocatalysts.

Oxidative enzymes are structurally and mechanistically diverse but a common feature of many is the use of molecular oxygen for catalysis. Enzymatic activation of oxygen is mediated by prosthetic groups/co-factors, most commonly flavins or metal ions coordinated by porphyrin or proteinaceous sulfur. A frequently observed byproduct of these reactions is hydrogen peroxide (H₂O₂). For example, P450 monooxygenases form H₂O₂ when substrate binding and electron transfer from NAD(P)H to the prosthetic group are not well coordinated. In the case of oxidases, molecular oxygen acts as a terminal electron acceptor and H₂O₂ forms along with the organic oxidation product in stoichiometric amounts. Because of its ubiquity, H₂O₂ formation can be considered a quasi-generic indicator of oxidative enzyme activity.

Enzymatic H₂O₂ formation is most commonly assayed by coupling of the H₂O₂-generating reaction to peroxidase-catalysed oxidation of fluorogenic or chromogenic indicators. In addition to considerable assay complexity requiring multiple liquid handling steps, the diffusive nature of the...
indicators makes them unsuitable for single-cell analyses and limits the plating density of screens executed on a solid support.

A more elegant means for H$_2$O$_2$ detection directly within living cells can be achieved by a genetically encoded probe termed HyPer.$^{19-21}$ HyPer makes use of permuted fluorophores (cpXFP) integrated into the regulatory domain of $E$. coli’s master regulator of oxidative stress, OxyR.$^{22}$ Oxidation of cysteine residues of the OxyR domain by H$_2$O$_2$ triggers a conformational change of cpXFP leading to a shift of the fluorescence spectrum that is indicative for cellular H$_2$O$_2$ levels. The HyPer sensor is especially suited for monitoring changes of the redox status of cells and cellular compartments but, due to the highly dynamic nature of the signal, is less suitable for enzyme screening applications where cumulative and irreversible reporters are preferred.

Here, we report the design and application of a genetically encoded transcription factor biosensor for detection of the intracellular H$_2$O$_2$ levels in $E$. coli developed specifically for enzyme screening applications. The suitability of the sensor for high-throughput enzyme screening was demonstrated with two $E$. coli whole-cell biocatalysts harbouring an alcohol oxidase or a cytochrome P450 monooxygenase, both of which generate H$_2$O$_2$ in the course of glycerol oxidation and toluene hydroxylation, respectively. We succeeded in rapidly isolating oxidase and monooxygenase variants with markedly increased activity towards non-natural substrates from large enzyme variant libraries, in screens on the level of colonies and single-cell levels using flow cytometry. The collected data demonstrates the broad scope, versatility, and compatibility of the presented H$_2$O$_2$-sensitive sensor with state-of-the-art high-throughput sampling protocols.

**Results and discussion**

**Biosensor development and characterization**

To non-invasively assess the oxidative activity of recombinant $E$. coli catalysts, we built an H$_2$O$_2$-sensitive genetically encoded sensor based on the master regulator protein of oxidative stress in $E$. coli, OxyR, and a cognate promoter, P$_{oxyS}$. At elevated intracellular H$_2$O$_2$ levels, OxyR is altered by oxidation of a cysteine residue (C199) and interacts with promoter P$_{oxyS}$ to modulate transcription.$^{24}$ We introduced the regulatory gene and the regulated promoter upstream of the genes encoding for green or red fluorescent proteins, GFP or RFP, to give plasmids pOxyR$_{GFP}$ and pOxyR$_{RFP}$ (Fig. 1A and Table S1†) that generate H$_2$O$_2$$_{RFP}$ or H$_2$O$_2$$_{GFP}$ fluorescence signals as outputs, respectively. As a pilot test, we supplied increasing amounts of H$_2$O$_2$ to $E$. coli BW25113(pOxyR$_{RFP}$) cells grown to mid-logarithmic phase and recorded H$_2$O$_2$$_{RFP}$ fluorescence change over time (Fig. S1A†). As H$_2$O$_2$ readily diffuses across the bacterial cell envelope,$^{25}$ H$_2$O$_2$$_{RFP}$ signal increased rapidly upon peroxide addition and reached a plateau within 180 minutes. The magnitude of the signal correlated well over an external concentration range of 5–100 µM H$_2$O$_2$ and non-linearly even up to 400 µM (Fig. 1A) which is consistent with both, high sensitivity and broad dynamic range of the sensor.

**Biosensor-mediated detection of oxidase activity**

Next, we investigated whether the OxyR sensor can be used to monitor intracellular H$_2$O$_2$ formation. For this purpose, we introduced plasmids pBAD$_{pcaoX}$ encoding variants of the glycerol oxidase from $Phanerochaete chrysosporium$ (PcAOX, EC 1.1.3.13)$^{26}$ into $E$. coli BW25113 ΔglpK/pOxyR$_{GFP}$. The resulting strain converts glycerol via the PcAOX enzyme as they no longer produce the housekeeping glycerol kinase GlpK.$^{27}$ Two PcAOX variants, wild type (WT) and variant F101S displaying 15-fold higher turnover frequency of glycerol,$^{28}$ were used to assess the H$_2$O$_2$$_{GFP}$ signal at increasing glycerol concentrations in a microtitre plate assay. After incubation with glycerol, cells expressing PcAOX F101S displayed the highest H$_2$O$_2$$_{GFP}$ signal which could be unambiguously distinguished from that of cells harbouring the WT enzyme. The H$_2$O$_2$$_{GFP}$ signal of PcAOX F101S-containing cells was on average 3.5-fold higher than that of cells harbouring the WT enzyme (Fig. S2A†) while the signal of the negative control (cells transformed with the same vector
indeed monitor the productive P450 hydroxylation reaction (in this example, o-cresol synthesis), a second genetically encoded sensor based on the transcription factor DmpR from Pseudomonas sp. \(^{27}\) was implemented (Fig. 1B). DmpR-based sensors have been previously used for screening of hydrolase\(^{28}\) and lyase\(^{29}\) mutant libraries but, to the best of our knowledge, not for monoxygenases. Given its high sensitivity and capacity to discriminate between tolualene and its phenolic derivatives,\(^{30}\) DmpR appeared suitable for detection of o-cresol formed upon hydroxylation of tolualene by cells that express the monoxygenase P450 BM3 from Bacillus megaterium (EC 1.14.14.1).\(^{31}\) We first increased the intracellular concentration of the DmpR regulator protein encoded on plasmid pDmpR_gfp by modifying the transcription of the dmpR gene via rational engineering of the P\(_f\) promoter (Fig. 1B and Table S2†) as reported elsewhere.\(^ {32}\) As a result, the o-cresol-triggered GFP signal (o-cresolGFP) in the presence of 10 \(\mu\)M o-cresol could be unambiguously measured within 60 minutes of incubation as compared to the >180 minutes required for the sensor employing the native P\(_f\) promoter (Fig. S1C†).

Next, the H\(_2\)O\(_2\)GFP and o-cresolGFP signals were used to simultaneously assess the formation of both enzymatic products. The plasmids encoding the two genetically encoded sensors were introduced into E. coli BL21 (DE3) lac\(_P\) harboring either P450 BM3 WT (encoded on pALXtreme-1a_p450_wt) or P450 BM3 M3 (on pALXtreme-1a_p450_m3). The two enzymes have been shown to exhibit very different H\(_2\)O\(_2\) and o-cresol formation rates with tolualene as substrate.\(^{33,34}\) Product formation was accurately reflected by o-cresolGFP and H\(_2\)O\(_2\)GFP signals measured on a single-cell level by flow cytometry (Fig. 2 and S3†). The cellular content of P450 BM3 WT and M3 monoxygenases were comparable (Fig. S6A†), which corroborates the notion that the observed differences in fluorescence are indeed resulting from the dissimilar catalytic performance of the two enzymes (Fig. 4C).

Taken together, the results illustrate that the proposed P450 monoxygenase activity determination method employing two biosensors allows to simultaneously monitor the intracellular production of hydroxylated organic products and H\(_2\)O\(_2\), and as such it has the potential to facilitate the isolation of improved enzyme variants.

**Biosensor-based screening of alcohol oxidase libraries**

As a first proof-of-principle screening application, we applied the OxyR sensor for pre-screening of a random mutagenesis library of PCAOX F101S aiming at enriching mutants with elevated glycerol-oxidizing capacity. Initially, we set out to measure H\(_2\)O\(_2\) formation as indicated by H\(_2\)O\(_2\)GFP signal using flow cytometry. By applying model libraries (mixtures of active and inactive E. coli strains at defined ratios), we could demonstrate efficient enrichment of the active strain during sorting (Fig. S2C†). However, treatment of E. coli cells with propidium iodide (PI, a DNA stain readily taken up only by damaged or dead cells\(^ {35}\)) indicated a correlation of H\(_2\)O\(_2\)GFP and PI-stain signals. This suggests that the extra H\(_2\)O\(_2\) generated as a result of the recombinant oxidase activity damages E. coli...
cells. It is also likely that those cells expressing the most active oxidase variants would be most severely affected. To mitigate this effect, we chose to perform the screening on agar plates. In addition to being simple to use ("plate, wait, and see") and easily parallelizable, this format allows maintaining high survival even for E. coli cells producing highly active oxidase variants (Fig. S4†). After incubating E. coli cells carrying mutagenized variants of the plasmid encoding pcaox gene on nutrient agar supplemented with 25 mM glycerol (20-fold lower than the reported Km of F101S), we visually inspected the plates and picked 90 clones displaying bright \( \text{H}_2\text{O}_2 \text{GFP} \) fluorescence. After their re-cultivation in liquid medium supplemented with glycerol, 85\% of these strains exhibited \( \text{H}_2\text{O}_2 \text{GFP} \) fluorescence between one and threefold the signal of the E. coli synthesizing the parent enzyme (±20\%) (Fig. 3A). This indicated that below 15\% of the picked E. coli strains were falsely assigned. As a control, 90 arbitrarily selected strains (colonies picked irrespective of the \( \text{H}_2\text{O}_2 \text{GFP} \) fluorescence) were also assayed. Less than 5\% of these strains displayed an \( \text{H}_2\text{O}_2 \text{GFP} \) signal similar to that of E. coli cells synthesizing the parental enzyme while the remaining ones were not distinguishable from the negative control (cells that do not express an oxidative enzyme). We could therefore efficiently enrich clones of elevated oxidative capacity. To verify this, we proceeded with the seven PcAOX F101S variants that triggered the strongest \( \text{H}_2\text{O}_2 \text{GFP} \) fluorescence, isolated the plasmids, and determined the pcaox gene sequence. Each of the variants carried a unique set of mutations (Table S2†). The PcAOX mutants were overproduced in an E. coli

Fig. 3 Screening of an error prone PCR library of PcAOX F101S. (A) Results from screening of an error-prone PCR library using the gene for PcAOX F101S as a template and plating of the transformed E. coli strains on LB agar supplemented with 100 mM glycerol. The relative activities of 90 highly \( \text{H}_2\text{O}_2 \text{GFP} \) fluorescent (light grey) and 90 arbitrarily selected (dark grey) PcAOX-variant producing E. coli strains are mapped; the relative activity (±20\%) of the parent is set to one and indicated by a dashed line. (B) Relative and absolute glycerol oxidation rates of hits identified in the initial screen in crude cell extract (upper panel) and with purified enzyme (lower panel).

Fig. 4 Screening of a saturation mutagenesis library of P450 BM3 using encoded sensors. (A) Flow cytometric pre-screening. The numbers added to the panels indicate the fraction (in \%) of the total events that fall into the \( \text{H}_2\text{O}_2 \text{RFP} \) (primary) and \( \text{o-cresolGFP} \) (secondary) gate. (B) Re-screening of individual isolates based on \( \text{o-cresolGFP} \) signal intensities after overnight incubation with substrate in a microtiter plate. (C) Simultaneous indirect whole-cell measurements of \( \text{o-cresol} \) and \( \text{H}_2\text{O}_2 \) formation by P450 BM3 WT and mutants M3 and M3.2 based on \( \text{H}_2\text{O}_2 \text{RFP} \) and \( \text{o-cresolGFP} \) signals. (D) Histogram plots of flow cytometric data recorded of a one-to-one mixture of cells expressing P450 BM3 wildtype and cells expressing the newly identified variant P450 BM3 M3.2. The two populations of cells can be discriminated relying on \( \text{H}_2\text{O}_2 \text{RFP} \) signal (top panel) but not on \( \text{o-cresolGFP} \) (bottom panel).

Top10 and oxidase activities in cell-free extracts [i.e., activities were normalized to total protein content in the lysate not to the cell-specific oxidase content thereby selecting for variants that were produced well and displayed in vitro activity] were determined using an ABTS/HRP-coupled assay (Fig. 3B and S3†). The three PcAOX variants that displayed the highest activity in lysate (D11, E8 and G12) were purified (Fig. S5A†) and their specific activity on glycerol was measured. All three variants showed improved initial activity on glycerol with variant displaying D11 displaying about 3-fold higher specific activity as compared to the parent enzyme, PcAOX F101S (Fig. 3B, bottom panel).

Biosensor-based screening of monooxygenase libraries

As a final proof-of-concept application, we set out to isolate more active monooxygenases variants for aromatic hydroxylation of toluene using the OxyR and DmpR sensor pair by
Table 1  Catalytic performance of P450 BM3 M3, M3.1 and M3.2 for aromatic hydroxylation of toluene

| P450 BM3 Mutations | Toluene conc. | NADPH oxidation | Coupling efficiency | o-Cresol formation |
|------------------|---------------|-----------------|---------------------|-------------------|
|                  | 1             | 10              |                     |                   |
| WT               | —             | 24 ± 1          | 10                  | 2                 |
| M3               | R48S Y51W A330F I401M | 89 ± 11         | 32 ± 2              | 29                |
| M3.1             | R48S Y51W E267Q A330F I401M | 652 ± 83      | 48 ± 3              | 313               |
| M3.2             | R48S Y51W A328P A330F I401M | 63 ± 6          | 19 ± 1              | 18                |
|                  | 10            | 45 ± 12         | 24 ± 11             | 11                |
|                  | 10            | 1066 ± 6        | 29 ± 3              | 309               |
|                  | 10            | 1609 ± 91       | 34 ± 1              | 547               |

The underlined mutations have been discovered in the course of this work. From Dennig et al., 2013. \(^a\) mM, \(^b\) \(\mu\)mol \(\mu\)mol \(^{-1}\) min \(^{-1}\), \(^c\) %, \(^d\) \(\mu\)mol \(\mu\)mol \(^{-1}\) min \(^{-1}\).

screening of a P450 BM3 M3 library. We performed biotransformation experiments with a pooled population of recombinant E. coli BL21 (DE3) lacP\(^T\) expressing a P450 BM3 M3 multi-site saturation mutagenesis library in liquid medium and simultaneously recorded \(\text{H}_2\text{O}_2\)^RFP and \(\text{o-cresol}^{\text{GFP}}\) signals at single-cell level by flow cytometry. Notably, low substrate load (0.1–0.2 mM toluene) and short incubation times (≤60 minutes) were applied to minimize unspecific background drift of \(\text{o-cresol}^{\text{GFP}}\) signal occurring at higher concentrations and/or longer incubation times (Fig. 4D). The reason for the deterioration of the \(\text{o-cresol}^{\text{GFP}}\) signal under these conditions was not fully elucidated. We suspect that because small phenolic compounds can rapidly penetrate bacterial membranes, the concentration of newly formed \(\text{o-cresol}\) steadily increased in the liquid medium to the point at which the highly-sensitive DmpR-based sensor was triggered in the entire E. coli population. Such a signal drift was significantly less pronounced for the \(\text{H}_2\text{O}_2\)^RFP signal. This is because \(\text{H}_2\text{O}_2\) can react with and oxidize cellular components (e.g. lipids, DNA, proteins, antioxidant molecules such as glutathione) and is actively degraded by several housekeeping enzymes (catalases and peroxidases) and only little \(\text{H}_2\text{O}_2\) presumably diffuses out of the cells. Accordingly, we sorted cells based on a high \(\text{H}_2\text{O}_2\)^RFP signal as a primary criterion and a high \(\text{o-cresol}^{\text{GFP}}\) signal as a secondary criterion. We argue that the combination and hierarchy of both signals provide means to lessen the effect associated with single-cell noise and background drift caused by membrane-permeable products (Fig. 4D). We spotted 384 gated E. coli cells on nutrient agar and obtained 42 colonies after overnight incubation (11% recovery). The \(\text{o-cresol}^{\text{GFP}}\) signal of individual E. coli strains in the presence of toluene was re-assessed in a microtiter plate assay and indicated that half of the clones displayed considerably lower signals in comparison to the parent (Fig. 4B). This result indicates a substantial degree of incorrect assignment likely due to the aforementioned background drift and noise inherent to single-cell measurements which could not be entirely prevented. The signal of the remainder of clones was in the range of the parent and one variant (M3.2) outstandingly displayed a 3-fold higher \(\text{o-cresol}^{\text{GFP}}\) signal than the parent. We selected M3.2 along with variant M3.1 featuring the second highest \(\text{o-cresol}^{\text{GFP}}\) fluorescence in re-screening assay and determined the DNA sequence of both genes. The enzymes were purified and characterized with respect to NADPH consumption (indicative of the overall \(\text{O}_2\) turnover frequency in the presence of toluene) and \(\text{o-cresol}\) formation (productive reaction). The \textit{in vitro} results (Table 1) recorded for M3.1 indicated 30% reduced activity to relative the parent at the substrate concentration (1 mM) used in the whole-cell re-screening assay. However, M3.2 displayed 11- and 1.7-fold increased product formation rates at low (1 mM) and high (10 mM) toluene, respectively. In addition, the degree of coupling of M3.2 was comparable to that of the parent at 1 mM and approximately 14% lower at 10 mM toluene. The \textit{in vivo} performance of this mutant and its parent measured with the genetically encoded sensors in a microtiter plate assay (Fig. 4C) correlated well with the \textit{in vitro} measurements. Taken together our results indicate that in a microtiter assay, the \(\text{H}_2\text{O}_2\)^RFP and \(\text{o-cresol}^{\text{GFP}}\) signals are suited for assignment of catalytic capacity of P450 BM3s within the complex environment provided by E. coli's cytosol and that the genetically encoded sensors can be employed with moderate success for high-throughput screening at a single cell level.

To demonstrate the crucial importance of the \(\text{H}_2\text{O}_2\)^RFP signal for flow cytometric sorting of P450 BM3 M3 libraries, a conversion of toluene by a one-to-one mixture of cells expressing P450 BM3 wildtype and the isolated variant P450 BM3 M3.2 was prepared and both \(\text{H}_2\text{O}_2\)^RFP and \(\text{o-cresol}^{\text{GFP}}\) signals were recorded over two hours (Fig. 4D). The result demonstrates that the two strains could be unambiguously distinguished based on the \(\text{H}_2\text{O}_2\)^RFP but not based on \(\text{o-cresol}^{\text{GFP}}\). This suggests that the \(\text{o-cresol}\) concentration in the culture broth gradually increased thereby activating the DmpR circuit and therefore \(\text{o-cresol}^{\text{GFP}}\) in all cells while the amount of secreted \(\text{H}_2\text{O}_2\) by cells expressing P450 BM3 M3.2 was not high enough to trigger a strong \(\text{H}_2\text{O}_2\)^RFP response in cells expressing the less active or inactive enzyme variants.

Conclusions

The genetically encoded sensors for high-throughput screening presented here enable reliable measurements of intracellular \(\text{H}_2\text{O}_2\) formed in the course of alcohol oxidase and P450 monoxygenase catalysed reactions formed as co-product or side product, respectively. \(\text{H}_2\text{O}_2\) has previously been shown to serve as an indicator for benchmarking of the performance of oxidative enzymes in enzyme-coupled assays. Classically, \(\text{H}_2\text{O}_2\),
can be assayed in the presence of a peroxidase (e.g. horse reddish peroxidase) which uses H\textsubscript{2}O\textsubscript{2} to oxidize dyes such as 2,2′-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid)\textsuperscript{18} or AmplexRed.\textsuperscript{17} The change of the spectral properties of the dyes can easily be recorded by plate readers or even by eye thereby allowing their application in a high-throughput setup. However, the supporting procedures include multiple manipulation steps for e.g. cell lysis and liquid handling, which complicates the workflow and limits the overall throughput.\textsuperscript{17,41}

Due to high degree of signal dynamics and a low signal-to-noise ratio barely exceeding an order of magnitude, the HyPer sensor design can only do so much for enzyme screening applications, in which integration times are short and the signal needs to be preserved for an extended period (in the range of hours) even if the reaction has been discontinued (e.g. to allow sample transfer or its pre-analytic storage). The OxyR sensor as reported here allows recording of H\textsubscript{2}O\textsubscript{2}-triggered signals over a wide dynamic range covering at least two orders of magnitude. When integrated directly into recombinant hosts expressing libraries of H\textsubscript{2}O\textsubscript{2}-liberating oxidative enzymes, the sensor proved compatible with several of the most frequently used platforms for high-throughput enzyme screening, i.e. plating on solid support followed by visual inspection of colonies, microtiter plate reader and flow cytometric measurements.

In summary, we present a method for activity assessment of oxidative enzymes directly in recombinant \textit{E. coli} expression hosts \textit{via} fluorescence measurements. The latter readouts are most frequently used for the execution of high-throughput screening protocols employed for biocatalyst optimization by directed evolution.\textsuperscript{42,43} The application of the genetically encoded OxyR sensor for enzyme screening provides advantages over existing approaches\textsuperscript{8} as it is economical (no need for chemical reagents), simpler to use (reduced number of manipulation steps, no cell lysis or compartmentalization required), and scalable (enabling ultra-high throughput). The OxyR sensor can therefore be readily used for activity screening of oxidizing biocatalysts provided that the substrates can be taken up by cells.

The high modularity of the OxyR sensor system ought to allow its application for detection of H\textsubscript{2}O\textsubscript{2} liberated in the in vitro characterisation of cytochrome P450 enzymes and ER contributed to oxidase engineering experiments. All authors discussed the results and commented on the manuscript.

Conflicts of interest
The authors declare the following competing financial interest(s): Elvira Romero is currently a Postdoctoral Researcher at AstraZeneca R&D Gothenburg.

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Data availability
Data are available within the article or its ESI materials.† The data that cannot be found in the article or in ESI materials† are available on request from the authors.

Author contributions
TK, MH and SP conceived the project; TK and MH designed the experiments; TK performed the majority of the experiments; TK and MH analysed the data and wrote the manuscript; AW performed in vitro characterisation of cytochrome P450 enzymes and ER contributed to oxidase engineering experiments. All authors discussed the results and commented on the manuscript.
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