LuxAB-Based Microbial Cell Factories for the Sensing, Manufacturing and Transformation of Industrial Aldehydes

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Abstract: The application of genetically encoded biosensors enables the detection of small molecules in living cells and has facilitated the characterization of enzymes, their directed evolution and the engineering of (natural) metabolic pathways. In this work, the LuxAB biosensor system from Photorhabdus luminescens was implemented in Escherichia coli to monitor the enzymatic production of aldehydes from primary alcohols and carboxylic acid substrates. A simple high-throughput assay utilized the bacterial luciferase—previously reported to only accept aliphatic long-chain aldehydes—to detect structurally diverse aldehydes, including aromatic and monoterpene aldehydes. LuxAB was used to screen the substrate scopes of three prokaryotic oxidoreductases: an alcohol dehydrogenase (Pseudomonas putida), a choline oxidase variant (Arthrobacter chlorophenolicus) and a carboxylic acid reductase (Mycobacterium marinum). Consequently, high-value aldehydes such as cinnamaldehyde, citral and citronellal could be produced in vivo in up to 80% yield. Furthermore, the dual role of LuxAB as sensor and monooxygenase, emitting bioluminescence through the oxidation of aldehydes to the corresponding carboxylates, promises implementation in artificial enzyme cascades for the synthesis of carboxylic acids. These findings advance the bio-based detection, preparation and transformation of industrially important aldehydes in living cells.

Keywords: biosensor; bioluminescence; luciferase; high-throughput screening; whole-cell biocatalysis; aldehyde production; enzyme cascade

1. Introduction

Great progress has been achieved in tailoring microorganisms for the biosynthesis of (non-)natural chemicals including but not restricted to: fatty acids and derivatives [1–3], aromatic and phenolic compounds [4–6] and secondary metabolites such as terpenoids [7,8]. In the context of the (bio-based) manufacturing of these compounds, aldehydes are of special interest not only as building blocks for pharmaceuticals but value-added products in the food, flavor and fragrance industries [9]. Enzymatically, aldehydes can be accessed through (1) decarboxylation reactions of 2-keto substrates [10] or fatty acids [11], (2) the readily established reductions of carboxylates by carboxylic acid reductases (CARs) [12–14] or (3) the oxidation of primary alcohols [9]. For the latter, only a few enzymes have been employed in living cells. One example is AlkJ, an alcohol dehydrogenase (ADH) from Pseudomonas putida (P. putida) being capable of oxidizing a variety of primary alcohols [15–17]. However, due to their high reactivity and cytotoxicity, aldehydes are rapidly metabolized by endogenous enzymes and do not accumulate in many microorganisms including (heterologous) hosts such as Escherichia coli (E. coli) [9,18,19]. To address this issue, the groups
of Atsumi and Prather constructed *E. coli* platform strains by targeted gene knock-outs that reduced the reduction of aliphatic and aromatic aldehydes to the corresponding primary alcohols [20,21]. The latter strain was termed *E. coli* RARE, exhibiting reduced aromatic aldehyde-reducing activity [21].

The functional expression of heterologous enzymes and the overall performance of microbial cell factories—not only for the synthesis of aldehydes—are usually assessed by the qualitative and quantitative analysis of target metabolites through chromatographic methods. Time-consuming sample preparation and low to moderate sample throughput impede the characterization of new biocatalysts as well as the development and optimization of microbial factories [22–24]. These bottlenecks have been addressed by the implementation of genetically encoded biosensor systems featuring (allosteric) transcription factors (TFs), riboswitches or enzyme-coupled devices to detect small molecules [25–29]. To date, biosensors have guided the directed evolution of enzymes [30–32], the engineering of (natural) metabolic pathways by the high-throughput (HT) detection of metabolites [33–35], and the dynamic regulation of genetic circuits to improve pathway performance, as well as other applications [3,28,36–41]. To sense aromatic aldehydes (e.g., benzaldehyde, cinnamaldehyde and vanillin), TF-based systems have been used [6,42,43]. Since TFs usually exhibit narrow ligand binding profiles, the detection of structurally different molecules is limited [4,42,44]. As a complementary tool, the luciferase LuxAB from *Photobacterium luminescens* (*P. luminescens*) was demonstrated to detect medium- and long-chain aliphatic aldehydes (C₈–C₁₄) [45]. LuxAB was also used to monitor fatty aldehyde/wax ester production [2] and non-native alkane synthesis pathways [25] in *Acinetobacter baylyi* ADPI (*A. baylyi*). As an alternative to biosensors, Ressmann et al. developed a HT assay to quantify different aldehydes in whole cells [46]: The heterologously produced aldehydes and exogenously added 2-amino-benzamidoximes (ABAOs) formed dihydroquinazolines, which could be detected spectrophotometrically [47]. While offering versatility, the assay depends on the multi-step synthesis of (substituted) ABAOs involving hazardous and toxic chemicals. Furthermore, the reaction with ABAOs irreversibly consumes target aldehydes.

In this work, the *P. luminescens* LuxAB biosensor, previously used in *A. baylyi*, was transferred to *E. coli*. This system provides a HT assay for aldehydes in vivo and was used to screen prokaryotic oxidoreductases for their aldehyde production capabilities. Since LuxAB is a monoxygenase (MO) that emits bioluminescence through the oxidation of aldehydes to the corresponding carboxylic acids (CAs) [45], it not only serves as a biosensor, but may also support the formation of industrially important carboxylates through an artificial enzymatic cascade starting from primary alcohols (Figure 1).

![Figure 1. LuxAB-based microbial cell factory. Aldehydes (b) can be produced from primary alcohols (a) or CAs (c) by heterologously expressed enzymes (cofactors omitted for clarity) and serve as substrates for the MO LuxAB, yielding the corresponding CAs and detectable bioluminescence. The conversion of (aromatic) aldehydes into the corresponding alcohols is decreased in the *E. coli* RARE strain [21]. The combination of the oxidation step (a,b) and LuxAB (highlighted in blue) represents a new cascade producing CAs.](image-url)
2. Results

2.1. LuxAB Biosensor Assembly and High-Troughput (HT) Detection of Aldehydes In Vivo

Aldehydes are greatly desired chemicals—for instance in the fragrance industry—and their bio-based production is highly demanded [9]. Hence, the fast characterization of new biocatalysts and easy detection of aldehydes are equally important. TF-based systems have been successfully employed to sense them but showed narrow ligand binding profiles [44]. This selectivity prompted us to implement the LuxAB system, described to sense a variety of aliphatic aldehydes [45], in the well-established biotechnology host E. coli.

For the construction of the biosensor device, the luxAB coding region was amplified from pAK400c/iluxAB_Cmr, which was previously constructed for the integration of luxAB into the A. baylyi genome and the monitoring of intracellular aldehyde levels (≤C18) [2,48]. Subcloning of luxAB under the control of a constitutive T5 promoter into the target pCDFDuet-1 vector was facilitated by a modified sequence and ligation-independent cloning (SLIC) procedure [49,50], yielding pLuxAB (Table S10). Functionality of the biosensor system was tested in E. coli RARE resting cells (RCs) as described in this study. Supplementation of octanal (2b), nonanal (3b) and decanal (4b) increased the bioluminescence by, roughly, 13,000-, 14,000- and 37,000-fold, respectively, in RCs expressing LuxAB in 96-well plates after 3 min (Figure 2A). The fold increase in bioluminescence above background caused by the presence of aldehydes was calculated as described in the materials and methods (Section 4.4). These results are supported by the initial research characterizing the substrate range of P. luminescens LuxAB [45].

Figure 2. Aldehyde detection by the LuxAB biosensor in E. coli. (A) the bioluminescence greatly increased in the presence of previously reported aliphatic aldehydes (2-4b). (B) selected new substrates of LuxAB included monoterpen aldehydes (15-17b) and aliphatic aldehydes with aromatic endgroups (19b and 22b). Experiments performed with RCs of E. coli RARE (OD_{600} ≈ 10.0) expressing LuxAB from pLuxAB in the presence of 1 mM aldehyde and 1% (v/v) ethanol as co-solvent; data presented as mean values + standard deviation (SD) of biological replicates (n ≥ 3).
To further investigate the aldehyde detection scope of LuxAB beyond reported aldehyde classes, the bioluminescence was monitored in the presence of different monoterpane (15–17b) and aromatic (18–25b) aldehydes at 1 mM (Figure 2B; selected examples) and 0.1 mM final concentration (Figure S5). Interestingly, all tested terpene aldehydes increased the bioluminescence, including citral (15b; applied as a commercial mixture of the E- and Z-isomers geraniol and nerol), citronellal (16b), with a clear preference for the (R)-enantiomer, and melonal (17b). (R)-16b and 17b, for example, increased the bioluminescence almost 2500- and 1500-fold, respectively. Surprisingly, 2-phenyl ethanol (19b) and 3-phenyl propanal (22b) also yielded bioluminescence (1700- and 230-fold increase, respectively, after 18 min; Figure 2B), whereas benzaldehyde (18b), cuminaldehyde (20b), trans-cinnamaldehyde (21b) and the tolualdehydes (23–25b) did not under these conditions (data not shown). The highest bioluminescence signals were detected at varying time points, which might indicate different rates and/or mechanisms of substrate uptake for aliphatic, monoterpane and aromatic aldehydes. Subsequently, the suitability of the LuxAB biosensor to detect aldehydes enzymatically produced in situ was determined. Therefore, three different prokaryotic oxidoreductases were selected: (1) the ADH AlkJ from P. putida, (2) a CAR from Mycobacterium marinum (CAR\textsubscript{Mm}) and (3) a choline oxidase variant from Arthrobacter chlorophenolicus (CO-6\textsubscript{Ac}), which has only been characterized in vitro [51]. AlkJ is known to oxidize aliphatic and aromatic primary alcohols [15–17], CAR\textsubscript{Mm} accepts both aliphatic and aromatic CAs [52–54] and CO-6\textsubscript{Ac} was initially engineered to oxidize hexanol (1a) [51].

The open reading frame (ORF) encoding alkJ was subcloned into a pACYC-derived backbone by SLIC, yielding pACYC\textsubscript{alkJ} as described in the materials and methods section (Section 4.2) and the Supplementary Materials. Similarly, the gene encoding a phosphopantetheinyl transferase from Nocardia iowensis (PPT\textsubscript{Ni}) and the target pACYC\textsubscript{Duet-1}/\textsubscript{car\textsubscript{Mm}} vector were assembled, yielding pACYC\textsubscript{Duet-1}/\textsubscript{car\textsubscript{Mm};pPT\textsubscript{Ni}}. PPTs are required to post-translationally modify apoCARs yielding catalytically active holoCAR enzymes [52,53]. For the expression of CO-6\textsubscript{Ac} in E. coli, the gene was codon-optimized, synthesized and subcloned into pET28a by the BioCat GmbH (Heidelberg, Germany). All constructs were designed to have compatible origins of replication for the stable maintenance with pLuxAB in the same cell as well as complementary antibiotic markers for the selection of co-transformants [55] (Table S10). Competent E. coli RARE cells were co-transformed with pLuxAB and either of the oxidoreductase-coding plasmids. Successful expression of enzymes was verified by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) analysis of whole-cell samples (Figure S4).

Finally, the HT conditions were employed to screen an extended substrate library containing the primary alcohols (1–7a, 18–25a), various diols (8–11a), simple monocarboxylates (1–7c, 18–25c), 6-hydroxy hexanoic acid (9c), 12-methyltridecanoic acid (12c), palmitoleic acid (13c), 12-hydroxy dodecanoic acid (14a) and dodecanedioic acid (14c).

All oxidoreductase-biosensor combinations yielded bioluminescence in the presence of substrates (Figure S3), suggesting the production of aldehydes in vivo. The HT screening indicated a broad substrate range for AlkJ and CAR\textsubscript{Mm} with an overlapping aldehyde production profile starting from primary alcohols and CAs, respectively. The highest bioluminescence could be observed with (unsubstituted) aliphatic substrates (1–7), also reflecting the preference of LuxAB to accept the corresponding aldehydes (Figure 2A). AlkJ and CAR\textsubscript{Mm} have been shown to produce 4b, for example, from decanol (4a) [56] and decanoic acid (4c) [54] and yielded 9281- and 989-fold increase in bioluminescence, respectively, after 1 h (Figure 3B). This almost 10-fold difference can be explained by the high bioluminescence background in RCs co-expressing LuxAB and CAR\textsubscript{Mm}: Endogenous (free) fatty acids can be reduced by the heterologous CAR enzyme. The corresponding fatty aldehydes serve as substrates for LuxAB, hence, yielding bioluminescence. Noteworthy, in all RCs co-expressing LuxAB and one of the oxidoreductases, the bioluminescence slightly increased over time in the presence of 1% (v/v) organic co-solvent and the absence of substrate (Figure S7). Therefore, the experimental cut-off (XCO) value was introduced...
to assess the fold increase in bioluminescence above background actually caused by the enzymatic production of aldehydes from substrates. The XCO was determined as described under materials and methods (Section 4.4) and in the Supplementary Materials.

**Figure 3. Cont.**
Figure 3. LuxAB-based HT screening of oxidoreductases in vivo. (A) bioluminescence suggests the production of aldehydes (b) from primary alcohols (a) by AlkJ (left) and CO-6Ac (middle) or CAs (c) by CAR_{Mm} (right; PPT_{Ni} for posttranslational modification of CAR_{Mm} omitted for clarity). The change in bioluminescence over time in response to the in situ production of aldehydes from (B) aliphatic substrates (1–14), (C) monoterpenes (15–16) and (D) aromatic compounds (18–25; continued on the next page). Shades of blue indicate the mean fold increase in bioluminescence based on biological replicates (n ≥ 3) above the XCO as defined in this study. Different scales reflect both varying background and the maximal bioluminescence. Experiments were performed in RCs of E. coli RARE (OD_{600} ≈ 10.0) co-expressing LuxAB and the described oxidoreductase in the presence of 1 mM substrate and 1% (v/v) ethanol as organic co-solvent; n.a. = not available (12–13a, 8c, 10–11c). LuxAB-based HT screening of oxidoreductases in vivo (continued). Change in bioluminescence over time in response to the in situ production of aldehydes by AlkJ (left) and CO-6Ac (middle) from primary aromatic alcohols (18–25a) or CAR_{Mm} (right) from aromatic CAs (18–25c).

The production of aldehydes from hexane-1,2-diol (8a), hexane-1,6-diol (9a), octane-1,2-diol (10a) and octane-1,8-diol (11a) by AlkJ was suggested by the rise in bioluminescence (e.g., largest fold increase for 8a of almost 2500; Figure 3B). To the best of our knowledge, these substrates have not been reported to be converted by AlkJ. The oxidase CO-6Ac had been shown to readily oxidize aliphatic 1–4a, the terminal diols 9a and 11a, the terpenoid alcohols geraniol (15a) and citronellol (16a), as well as the aromatics benzyl alcohol (18a), 2-phenylethanol (19a) and cinnamyl alcohol (21a) in vitro [51]. The same substrate scope was observed under the HT assay conditions in vivo (Figure 3B–D). The low maximal bioluminescence in these reactions is probably caused by the competition for (dissolved) O₂ by LuxAB and CO-6Ac, and the cytotoxic H₂O₂ produced by the oxidase (Figure 3A) [57], decreasing cell viability and indirectly quenching bioluminescence. Additionally, the availability of flavins in E. coli might present another bottleneck [58], as discussed below. As mentioned above, the direct addition of aromatic aldehydes 18b and 21b, for example, did not increase the bioluminescence in RCs solely expressing LuxAB. This behavior can be explained by the cytotoxicity of aldehydes at elevated concentrations [9]. Nonetheless, the formation of aldehydes in situ by AlkJ, CO-6Ac and CAR_{Mm} seemed compatible with the LuxAB biosensor system based on the detection of bioluminescence in samples presumably containing 15–22b (Figure 3D). Furthermore, the production of the value-added monoterpene aldehydes 15–16b was suggested (Figure 3C); 15–16b have not been reported as products of AlkJ- or CAR-catalyzed reactions. Consequently, biotransformations in whole cells expressing AlkJ or CAR_{Mm}/PPT_{Ni} were performed to confirm the production of selected aldehydes, based on the results of the HT assay.

2.2. Microbial Cell Factories for the Production and Transformation of Value-Added Aldehydes

Biotransformations were carried out in RCs of E. coli RARE transformants. Focus was on unreported substrates including 4-isopropylbenzyl alcohol (20a), cinnamyl alcohol...
(21a), 2-methylbenzyl alcohol (23a) and 4-methylbenzyl alcohol (25a) for AlkJ (Figure 4) and the corresponding terpenoid substrates (15–16) for both AlkJ and CARMm (Figure 5A).

**Figure 4.** Extended scope of AlkJ towards aromatic compounds. Oxidation of the new alcohol substrates 20–21a, 23a and 25a to the corresponding aldehydes (b) by AlkJ and overoxidation to CAs (c) by endogenous enzymes. In biotransformations of 21a, 3-phenyl propanol (22a) was detected as byproduct. Experiments were performed in RCs of *E. coli* RARE (OD_{600} ≈ 10.0) expressing AlkJ from pACYQ in the presence of 5 mM alcohol (a) and 5% (v/v) ethanol as co-solvent. Sampling: 0 h (after the addition of substrate and mixing), 1 h and 24 h. Reduced recoveries attributed to low solubility in resting cell medium (RCM) and/or volatility of compounds; 100% recovery represents the complete retrieval of the amount of substance added. GC yields presented as mean values + SD (%) of biological replicates (n = 3).

GC/FID analysis confirmed the AlkJ-catalyzed production of 20–21b and 25b from the corresponding aromatic alcohols with the highest GC yield for the industrially important 21b (56.5 ± 6.8% after 24 h). Besides the overoxidized cinnamic acid (21c; 8.5 ± 2.2%), 3-phenyl propanol (22a; 6.5 ± 4.2%)—lacking the C = C bond in the side chain—could be detected; both are expected byproducts formed by endogenous host enzymes [15,59,60]. Further, the tested aromatic alcohols with aliphatic substituents in the para-position were utilized by AlkJ; 20b and 4-methylbenzaldehyde (25b) were readily produced, whereas the ortho-substituted 23a only showed a low level of conversion (Figure 4). The same preference for aromatic aldehydes with aliphatic substituents in the para-position is indicated for LuxAB, based on the HT assay results (Figure 3D). This prediction was substantiated by biotransformations in RCs of *E. coli* RARE expressing CARMm/PPT_{Ni}: The ortho-, meta- and para-substituted 23–25c were reduced to 2-methylbenzaldehyde (23b), 3-methylbenzaldehyde (24b) and 25b, respectively (Figure S8). Since 23b and 24b are poor substrates for LuxAB, they could not be unambiguously detected in the HT assay (Figure 3D).
Production of (monoterpene) aldehydes by CAR

GC/FID analysis also confirmed the enzymatic production of value-added monoterpene aldehydes in vivo. CARMm efficiently converted geranic acid (15c) and the two enantiomers of citronelic acid—(R)- and (S)-16c—to the corresponding aldehydes. The reduction of 15c yielded 49.0 ± 2.7% of the desired 15b and the over-reduced 15a (22.9 ± 1.1%). Since E. coli RARE was engineered to specifically increase the persistence of aromatic aldehydes, the reduction of other aldehyde classes can be expected [21]. Furthermore, 16b, which lacks the proximal C=C double bond, was detected (11.0 ± 0.8%) after 24 h (Figure 5). As before, these byproducts are produced by host enzymatic activities [9,15,20,59–61]. Similarly, the reductions of (R)- and (S)-16c yielded (R)-16b (76.1 ± 2.2%) and (S)-16b (63.9 ± 10.3%), as well as the over-reduced 16a and the over-oxidized 15c (Figure 5; see also: Table S11). Again, these results substantiated the extended aldehyde detection scope of LuxAB (Figure 3C). AlkJ showed low activity towards (R)- and (S)-16a, yielding 11.8 ± 1.2% (R)-16b and 2.4 ± 0.2% (S)-16b, respectively. Interestingly, only 15c—not 16c—was detected as the byproduct in low amounts (Figure 6A; Table S12). Under the same conditions, AlkJ efficiently oxidized 15a, exclusively yielding 83.8 ± 1.0% of the important aroma compound 15b after 24 h; the overoxidized 15c was not detected (Figure 6A).

Like the flavin adenine dinucleotide (FAD)-dependent ADH AlkJ, potential endogenous...
EREDs such as N-ethylmaleimide reductase [62] or 2,4-dienoyl-CoA reductase [63] are flavoproteins. That the reduction of the proximal double bond was not observed in biotransformations of 15a could be explained by the competition for FAD and/or the depletion of the precursor flavin mononucleotide (FMN) [58]. This bottleneck might also be the cause for the only slightly increased amounts of CAs in biotransformations employing RCs co-expressing AlkJ and the FMN-dependent MO LuxAB (Figure 6; see also Figure 1). This does not necessarily contradict the high bioluminescence signals of AlkJ/LuxAB under HT assay conditions since luciferases are very good photoemitters in terms of quantum yield and the detection is highly sensitive [64]. Starting from 4a, (R)-16a or (S)-16a, the amounts of the observed overoxidation products increased 1.3-fold (4c), 1.4-fold (15c) and 2.1-fold (15c), respectively, in cells co-expressing AlkJ and LuxAB; 15c could be detected in traces in biotransformations starting from 15a (Figure 6B), which was not observed in reactions employing AlkJ alone (Figure 6A). Although subtle, these findings suggest a new enzymatic cascade transforming alcohols into carboxylates via two steps in living cells.

**Figure 6.** Production of (monoterpene) aldehydes by AlkJ and transformation by LuxAB in vivo. Oxidation of 15–16a to the desired aldehydes (b) and further oxidation to the corresponding CAs (c) by (A) endogenous *E. coli* enzymes and (B) LuxAB. The oxidation of 4a was used as positive control and yielded byproducts originating from pathways for the metabolism of fatty acids [65,66]. The exact compositions including byproducts are given in Table S12. Experiments were performed in RCs of *E. coli* RARE (OD<sub>600</sub> ≈ 10.0) expressing (A) only AlkJ or (B) AlkJ and LuxAB from pAYCQ and pLuxAB, respectively, in the presence of 5 mM alcohol (a) and 5% (v/v) ethanol as organic co-solvent. Sampling: 0 h (after the addition of substrate and mixing), 1 h and 24 h; reduced recoveries attributed to low solubility in RCM, volatility and/or metabolization of compounds. GC yields presented as mean values + SD (%) of biological replicates (n ≥ 2).

### 3. Discussion and Conclusions

The continuously expanding number of new biocatalysts from both natural resources and protein engineering endeavors demands for tools for their rapid characterization [67,68]. In this context, genetically encoded biosensor systems have been embraced since they translate the detection of small molecules into a measurable output such as fluorescence or bioluminescence [34,69].

In this study, the LuxAB system from *P. luminescens*, emitting bioluminescence through the oxidation of aldehydes to the corresponding CAs [45], was successfully implemented in *E. coli* RARE (Figure 1). This effort led to the establishment of a simple HT assay...
that revealed the capability of LuxAB to detect structurally diverse aldehydes, including monoterpene and aromatic aldehydes in living cells (Figure 2). The versatility of the assay was demonstrated by the screening of three different microbial oxidoreductases—AlkJ, CO-6Ac and CARMm—against an extended substrate library, producing up to 25 industrially relevant aldehydes (Figure 3). Under HT screening conditions, both the background bioluminescence and the maximal bioluminescence signal depended on the expressed oxidoreductase. AlkJ/LuxAB yielded the highest maximal-fold increase in bioluminescence in the presence of the preferred aliphatic substrates (e.g., >9000-fold for 4a). In comparison, LuxAB/CARMm and LuxAB/CO-6Ac only increased the maximal bioluminescence >900-fold (4c) and ≈ 100-fold (3a), respectively (Figure 3B). These deviations are not solely based on the substrate preferences of the target oxidoreductase but can be explained by their distinct enzymatic properties including the acceptance of intracellular fatty acids as substrates by CARMm, increasing the background bioluminescence or the production of H2O2 by the mutant choline oxidase CO-6Ac, resulting in decreased cellular viability. The introduction of an XCO value sufficed to address varying backgrounds and allowed the confirmation of the substrate ranges of AlkJ, CARMm and CO-6Ac in vivo. The latter only had been assessed before in vitro [51]. Furthermore, new substrates for the ADH AlkJ and CARMm could be identified and confirmed in whole-cell transformations and subsequent GC/FID analysis. Biotransformations also revealed the limits of the detection scope of the LuxAB biosensor system: Although the aromatic aldehydes 23–24b were readily produced by CARMm (Figure S8), for example, they are poor substrates for LuxAB and could not be efficiently detected under HT assay conditions; 23b was also produced from 23a by AlkJ previously [15]. On the other hand, poor substrates for a given oxidoreductase yielding low amounts of aldehydes that are well-accepted by LuxAB might emit high bioluminescence, consequently, suggesting a deceptive oxidoreductase scope.

In addition to 20b, 23b and 25b, AlkJ yielded the flavorant cinnamaldehyde (21b) in satisfying yields. In a cascade reaction catalyzed by endogenous host enzymes, 20b, 23b and 25b were oxidized to the corresponding CAs (Figure 4). CARMm efficiently reduced the newly identified carboxylate substrates (R)- and (S)-16c to the corresponding monoterpene aldehydes in 76.1 ± 2.2% and 63.9 ± 10.3% GC yields, respectively (Figure 5). Citronellal (16b) is not only used as a scent, but has insect repellent and antifungal properties [70,71]. Excitingly, the AlkJ-catalyzed oxidation of 15a exclusively formed 15b (>80% GC yield; Figure 6A), offering a biocatalytic alternative to chemical oxidation procedures [72] and enabling combinations with other enzymatic transformations, for example, the XenA-mediated reduction of the proximal C = C double bond, yielding 16b, as suggested by Richardson and co-workers [61]. Citral (15b) is widely used in perfumes, as mosquito repellent [73] and is an important precursor for other terpenes [74] and carotenoids including vitamin A [75]. Albeit low, this study also confirmed the MO activity of LuxAB in a cascade set-up with AlkJ, indicated by the detectable increase in carboxylate products (Figure 6B).

In summary, the genetically-encoded LuxAB proved to be a versatile biosensor for the HT detection of structurally diverse aldehyde compounds, without the need of undesirable auxiliary chemicals such as ABAOs. Additionally, the fast detection of aldehydes by the enzyme-based biosensor is advantageous to TF-based systems. The applicability of the HT assay was demonstrated by the extension of the substrate scopes of different oxidoreductases and the bio-based production of monoterpene aldehydes, amongst others. These findings will facilitate the identification and characterization of novel biocatalysts in the future, advance the enzymatic preparation of industrial aldehydes and point towards their diversification through (LuxAB-based) microbial cell factories.

4. Materials and Methods

4.1. General Information

Chemicals, reagents and reference compounds were purchased from Sigma-Aldrich (Buchs, Switzerland), Roth (Karlsruhe, Germany) or TCI Chemicals (Tokyo, Japan) and used without further purification, unless stated otherwise. Table S13 provides an overview
of all compounds (1–25) analyzed in this study. Solvents (GC grade) were purchased from Sigma-Aldrich or J.T. Baker (Schwerte, Germany).

The following E. coli laboratory strains were purchased from Thermo Scientific/Invitrogen (Darmstadt, Germany): BL21 (DE3), DH5α and TOP10. E. coli RARE was acquired from the Prather group [21].

Unless stated otherwise, chemically competent E. coli cells were produced by well-established protocols using CaCl$_2$ (0.1 M) and transformed with plasmid DNA (25–100 ng·µL$^{-1}$) by heat-shock at 42°C for 45 s (see: Supplementary Materials). Recovery was performed in SOC medium (2% tryptone, 0.5% yeast extract, 10 mM NaCl, 2.5 mM KCl, 10 mM MgCl$_2$, 10 mM MgSO$_4$ and 20 mM glucose) at 37°C with vigorous shaking. For efficient transformation of E. coli RARE, plasmids were passed through E. coli DH5α before. Selection of transformants was performed in lysogeny broth (LB) medium (25 g·L$^{-1}$; Sigma-Aldrich) or on LB agar plates containing 1.5% agar-agar (Roth) in the presence of the appropriate antibiotic(s) (final concentration): chloramphenicol (34 µg·mL$^{-1}$), kanamycin (50 µg·mL$^{-1}$) and streptomycin (25 µg·mL$^{-1}$). Only half the concentration of antibiotics was used for the selection and subsequent cultivation of strains harboring two plasmids. Co-transformants were produced by transforming chemically competent E. coli RARE harboring pLuxAB with the desired oxidoreductase-coding plasmid as described above.

4.2. Plasmid Assemblies by Sequence- and Ligation-Independent Cloning (SLIC) Techniques

Standard DNA oligonucleotides (desalted) were ordered from Thermo Fisher Scientific/Invitrogen and dissolved in nuclease-free water (Invitrogen). The resulting stock solutions (100 µM) were further diluted and used as primers for PCRs (5 µM) or Sanger sequencing (10 µM). PCRs were performed in a thermal cycler (Biometra TAdvanced, Analytik Jena, Jena, Germany) with Pfu$^+$ or OptiTaq DNA polymerase (Roboklon, Berlin, Germany) or Q5® high-fidelity (Q5®-HF) DNA polymerase (New England Biolabs, Frankfurt/Main, Germany); premixed dNTPS (10 mM each) were purchased from Roth. Other enzymes for the manipulation of DNA were purchased from Thermo Fisher Scientific or New England Biolabs.

PCR products were purified with QIAquick PCR & Gel Cleanup Kit (QIAGEN, Hilden, Germany) and plasmid DNA isolated with innuPREP Plasmid Mini Kit 2.0 (Analytik Jena) as instructed by the suppliers. DNA concentration was determined by NanoDrop™2000 spectrophotometer (Thermo Fisher Scientific, Darmstadt, Germany). Sanger sequencing was performed by Eurofins Genomics (Ebersberg, Germany).

In this study, plasmids were assembled from two linear DNA fragments via homologous overhangs attached by PCR, following adapted protocols from Li et al. [76] or Wiesinger et al. [50]. The latter employed a seamless and ligation-independent cloning extract (SLiCE) that was prepared according to Zhang et al. from E. coli TOP10 cultures [49]. Primer sequences and templates are given in Tables S9 and S10, respectively, in the Supplementary Materials. Optimal annealing temperatures ($T_a$) were determined by gradient PCR (45–65 °C), unless stated otherwise. Composition of PCR mixtures and construct-specific thermal cycle conditions are given in the Supplementary Materials.

For the construction of pCDFduo/luxAB, herein referred to as pLuxAB, the DNA fragment encoding the luxAB subunits was amplified from pAK400c/iluxAB_Cm$^r$ [48] with the primer pair LuxA_T5 F/LuxB R, which introduced flanking 15 bp-overhangs complementary to the target pCDFDuet-1 vector for the subsequent assembly, and Pfu$^+$ ($T_a = 45.0$ °C; Figure S1). The pCDF backbone was amplified with primers pCDF F/R and OptiTaq ($T_a = 48.0$ °C as calculated from the melting temperature of primers; Figure S1). The correct assembly by SLiCE was confirmed by Sanger sequencing of isolated plasmid DNA from single colonies of E. coli BL21 (DE3) transformants selected on LB agar plates supplemented with streptomycin.

For the construction of pAYCQ/alkJ, the DNA fragment harboring the alkJ fragment was amplified from pGEc47 [77] with the primer pair AlkJ F/R, introducing 15 bp-overhangs complementary to the target pACYC vector, and Pfu$^+$ ($T_a = 48.6$ °C; Figure S2). The pACYC
backbone was amplified with the primers pACYC F/pACYC-2 R and OptiTaq (T_a = 48.6 °C; Figure S2). Purified DNA fragments were processed and incubated with the SLiCE. Subsequently, chemically competent E. coli TOP10 cells were transformed with the assembly mixtures. Colonies were selected on LB agar plates containing chloramphenicol the next day. Sanger sequencing of isolated plasmid DNA revealed a truncated ORF of the alk/j gene in all samples (see also: Table S10). The ORF was restored using the Q5® Site-Directed Mutagenesis Kit (New England Biolabs) with the primer pair AJQ5 F/R (T_a = 61.0 °C as calculated by the NEBaseChange™ tool available from: http://nebasechanger.neb.com/ (accessed on 8 September 2019) and following the instructions of the supplier. Finally, the completeness of the alk/j ORF was confirmed by Sanger sequencing.

For the construction of pACYCDuet-1/car_Mm:ppt_Ni, chemically competent E. coli TOP10 cells were directly transformed with the assembly mixture (molar ratio of insert to backbone of 3:1) according to Li et al. [76] after DpnI digestion. Therefore, the ppt_Ni insert had been amplified from pCDF/ppt_Ni with the primer pair Npt F/R and Q5®-HF (T_a = 57.0 °C; Figure S3). The pACYCDuet-1 backbone harboring the car_Mm gene was amplified with the primers pCAR_Npt F/R and Q5®-HF (T_a = 64.0 °C; Figure S3). The correct assembly was confirmed by Sanger sequencing of isolated plasmid DNA from single colonies of transformants selected on LB agar plates supplemented with chloramphenicol.

The gene encoding the choline oxidase variant was codon-optimized for the expression in E. coli, synthesized and subcloned into pET28a by the BioCat GmbH (Heidelberg, Germany).

4.3. Standard Conditions for Enzyme Production and the Preparation of Resting Cells (RCs)

Protein production (AlkJ, CAR_Mm / PPT_Ni, CO-ßAc, and LuxAB) was performed in E. coli RARE transformants cultivated in auto-induction medium (AIM; 2.5% LB medium, 1 mM MgSO_4, 25 mM (NH_4)_2SO_4, 50 mM KH_2PO_4, 50 mM Na_2HPO_4, 5% glycerol, 0.5% glucose and 2% α-lactose) adapted from Studier [78]. Briefly, a single colony of the desired strain was grown in LB medium containing the appropriate antibiotic(s) at 37 °C (180 rpm) for 12–16 h. AIM supplemented with antibiotic(s) was inoculated with 0.2% (v/v) preculture in baffled flasks and incubated in Infors HT Multitron incubator shakers (Bottmingen, Switzerland) at 37 °C (180 rpm) for 4–6 h (6 h for co-transformants, 5 h for pLuxAB transformants and 4 h for all others). Enzyme production was performed at 20 °C (150 rpm) for 16–20 h. The optical density at 600 nm (OD_600) of cultures was determined with a UV-1280 spectrophotometer (Shimadzu, Kyoto, Japan). Cells were harvested by centrifugation (6000 x g, 4 °C) for 20 min using a Heraeus Fresco 17 centrifuge or a Heraeus Labofuge 400R (Thermo Fisher Scientific). The cell pellet was resuspended in RCM (22 mM KH_2PO_4, 42 mM Na_2HPO_4, 8.56 mM NaCl, 1 mM MgSO_4, 1 mM CaCl_2 and 1% glucose) until an OD_600 ≈ 10.0 was reached. RCs were used on the day of preparation.

Protein expression was confirmed by 12.5% (v/v) SDS-PAGE analysis of whole-cell samples normalized to OD_600 = 7.0, using the Mini-PROTEAN electrophoresis system (Bio-Rad, Feldkirchen, Germany) and following standard protocols (e.g., [50]). Gels were stained with InstantBlue™ Protein Stain (Expedeon, Heidelberg, Germany).

4.4. LuxAB-Based HT Screening of Enzymes In Vivo

RCs co-expressing LuxAB and the enzyme of interest were prepared as described above. To 198 µL, RCs (OD_600 ≈ 10.0) per well, 2 µL stock solution of the target substrate (0.1 M in dimethyl sulfoxide or ethanol) were added to a final concentration of 1 mM substrate (V_0 = 200 µL containing 1% (v/v)) organic co-solvent) in 96-well plates (flat bottom, black polystyrene [50,65,76]; Greiner Bio-One, Frickenhausen, Germany). It was mixed gently and the bioluminescence measured immediately on a Varioskan™ LUX multimode plate reader (Thermo Fisher Scientific). The change in bioluminescence was followed at 25 °C for up to 1 h.

To assess the fold increase in bioluminescence above background (b_x) caused by the enzymatic production of aldehydes from substrates, the blank-corrected bioluminescence signal at the corresponding time point (t_x) was divided by the bioluminescence signal
before substrate addition (t₀; see also: Supplementary Materials). The blank value is referred to the bioluminescence in RC suspensions before the addition of any organic compound. In parallel, the increase in bioluminescence in the presence of 1% (v/v) organic co-solvent in RCs expressing LuxAB (or in combination with the oxidoreductase of interest) was monitored and the fold increase calculated as before, yielding the cut-off value. Substrate-enzyme combinations exhibiting bioluminescence greater than the cut-off value + SD, referred to as the XCO (see also: Figure S7), were re-screened in whole-cell biotransformations expressing the corresponding oxidoreductase alone.

4.5. Whole-Cell Biotransformations and Gas Chromatographic (GC) Analysis

RCs (OD₆₀₀ ≈ 10.0) expressing the oxidoreductase of interest or expressing LuxAB and AlkJ for cascade reactions were prepared as described under Section 4.3; whole-cell biotransformations were performed in glass vials with screw-caps (4 mL) at 5 mM substrate concentration for alcohols and carboxylates in the presence of 5% (v/v) organic co-solvent (Vtotal = 0.5 mL) in Infors HT Multitron incubator shakers at 25 °C (220–250 rpm) for 0–24 h. For GC analysis, samples (100 µL) of the biotransformation mixtures were taken immediately after the addition of substrate and mixing (t ≈ 0 h), 1 h and 24 h. Unless for biotransformations involving monoterpenoid aldehydes (e.g., 15b) and related compounds [74], samples were acidified with 2 M HCl (10 µL) and extracted two times with ethyl acetate (200 µL) containing 1 mM methyl benzoate as internal standard (IS) by vortexing for 30–45 s. It was centrifuged (13,000 × g, 4 °C) for 1 min. The combined organic phases were dried over Na₂SO₄ and transferred into a GC vial with insert, capped and submitted to GC analysis. Compound identification was performed by the comparisons of retention times of commercial standards (Table S13), unless stated otherwise; quantification and calculation of GC yields were performed by standard calibrations or using relative response factors (see: Supplementary Materials).

GC analysis (GC-2010 Plus, Shimadzu) using a flame ionization detector (FID; Shimadzu) was performed on a ZB-5MSi column (length: 30 m; inner diameter: 0.25 mm; film thickness: 0.25 µm) from Phenomenex (Torrance, USA). GC/FID method (hydrogen, 0.96 mL min⁻¹ flow rate; injector and detector: 300 °C): 100 °C, hold 1 min, 20 °C per min to 250 °C, hold 5 min; total time: 13.5 min. GC/MS analysis (GCMS-QP2010 SE, Shimadzu) was performed on the same column. GC/MS method (helium, 1.00 mL min⁻¹ flow rate; injector: 280 °C, ion source and interface: 260 °C): 100 °C, hold 5 min, 20 °C per min to 250 °C, hold 5 min, 20 °C per min to 280 °C and hold 5 min; total time: 24.0 min.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/catal11080953/s1; Figure S1: DNA fragments for the assembly of pLuxAB, Figure S2: DNA fragments for the assembly of pACYC/alkJtrnc, Figure S3: DNA fragments for the assembly of pACYCDuet-1/carMm:pptNi, Figure S4: SDS-PAGE analysis of whole-cell samples, Figure S5: LuxAB-based detection of aldehydes at low concentration in E. coli, Figure S6: LuxAB-based detection of aldehydes in E. coli (background and negative controls), Figure S7: Background luminescence and determination of the XCO value, Figure S8: Enzymatic reduction of toluic acids by CARMm; Table S1: PCR mixtures for the assembly of pLuxAB, Table S2: Optimized thermal cycle conditions for the assembly of pLuxAB, Table S3: PCR mixtures for the assembly of pACYC/alkJ, Table S4: Optimized thermal cycle conditions for the assembly of pACYC/alkJ, Table S5: Q5® mutagenesis reaction mixture, Table S6: Thermal cycle conditions for Q5® mutagenesis, Table S7: PCR mixtures for the assembly of pACYCDuet-1/carMm:pptNi, Table S8: Optimized thermal cycle conditions for the assembly of pACYCDuet-1/carMm:pptNi, Table S9: List of DNA oligonucleotides used in this study, Table S10: List of strains and plasmids used in this study, Table S11: Production of (monoterpenic) aldehydes by CARMm and transformation in vivo, Table S12: Production of (monoterpenic) aldehydes by AlkJ and transformation by LuxAB, Table S13: List of compounds analyzed in this study.
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