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- This study introduces a new sustainable aquacatalytic system utilizing medium-chain fatty acids (MCFAs), especially caprylic acid, for Brønsted acid-catalyzed carbonyl allylation under "on-water" conditions. The system operates efficiently using seawater or edible sea salt, offering a low-cost and environmentally benign alternative to conventional solvent condition.
- 2. The core innovation exploits the salting-out effect to suppress micelle formation and enhance interfacial reactivity, enabling catalyst-free aldehyde allylation and caprylic acid-catalyzed ketone allylation in aqueous media. The reaction conditions are mild, scalable, and compatible with DNA-tagged substrates, opening avenues for pharmaceutical synthesis and drug discovery.
- 3. Future directions may involve broadening the scope of MCFA-based catalysis across other aqueous transformations, integrating renewable MCFA sources from food or waste streams, and developing entirely water-based catalytic platforms to further enhance the system's green chemistry potential.

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View Article Online DOI: 10.1039/D5GC01441G

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Received 00th January 20xx, Accepted 00th January 20xx

DOI: 10.1039/x0xx00000x

"On-Seawater" Accelerated Aquacatalysis by Edible Fatty Acid: Harnessing the Remarkable Salting-Out Effect

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Medium-chain fatty acids (MCFAs) are widely recognized for their metabolic and therapeutic benefits, yet their potential as catalysts for chemical reactions remains elusive. Herein, we report the development of a sustainable aquacatalytic system utilizing MCFAs, particularly caprylic acid, for Brønsted acid-catalyzed carbonyl allylation under "on-water" conditions. This approach leveraged the salting-out effect induced by NaCl to suppress micelle formation and enhance interfacial catalysis. By employing unrefined seawater and edible sea salt as cost-effective additives, the system enables the efficient allylation of aldehydes without a catalyst and the caprylic acid-catalyzed allylation of ketones. This reaction is scalable to gram-scale synthesis using a chromatography-free purification process, offering a practical and sustainable route for producing active pharmaceutical ingredients. Furthermore, the mild reaction conditions and compatibility with aqueous media facilitated successful on-DNA allylation, underscoring the potential of DNA-encoded library applications in drug discovery. This study highlights the unprecedented utility of MCFAs as renewable catalysts and establishes a versatile and environmentally friendly platform for aquatic organic transformations.

Introduction

Medium-chain fatty acids (MCFAs), which are carboxylic acids with carbon chain lengths of 6-12, are rapidly digested and absorbed by the human body and serve as an efficient energy source¹⁻⁴. In addition to their metabolic roles, MCFAs offer a range of health benefits, including weight loss support⁵⁻⁷, reduction in cardiovascular disease risk⁸, enhancement of dietinduced thermogenesis and satiety⁹, antimicrobial activity¹⁰, and improved digestive efficiency¹¹. These unique properties have garnered significant attention in nutritional science and medical research, as they are valuable components in functional foods and therapeutic interventions^{3,12–14}. In addition to their biological significance, MCFAs exhibit distinctive amphiphilic properties owing to their chemical structure, which features a hydrophilic head and a hydrophobic tail. This structure imparts excellent surfactant-like characteristics, including biocompatibility, biodegradability, low toxicity, and high specificity. These characteristics make MCFAs suitable for applications such as drug delivery systems in both synthetic and biomedical contexts¹⁵. Notably, in aqueous environments, MCFAs can self-assemble into micelles at their critical micelle concentration (CMC), which enables them to perform stably and efficiently under physiological conditions^{16,17}. Caprylic acid

(C8:0), a prominent MCFA, exemplifies the advantages of this class of compound. The carbon chain enhances its solubility and metabolic efficiency, facilitating its rapid absorption and direct utilization as an energy source in the liver¹⁸. In addition, its potent antimicrobial activity against various pathogens underscores its utility in functional foods and therapeutic applications¹⁰. Furthermore, their ability to form stable micelles highlights their potential as versatile carriers^{19,20}. However, their role as catalysts in chemical reactions remains entirely elusive (Fig. 1A).

Recent studies by our group have revealed that various catalytic transformations can be effectively accelerated under "onwater" conditions interface²¹⁻²⁵, where reactions occur at the interface between bulk-water and compact-oil phases^{26,27}. Mechanisms such as Brønsted superacid²¹, Brønsted superbase^{22,24}, and photocatalysis²⁵ were observed, which significantly augmented the catalytic activity at the interface²⁸. This phenomenon is mainly observed in addition-type reactions (e.g., Michael addition and [2+2] cycloaddition), which typically involve a decrease in the volume of the transition state²⁹. Although multiple factors may contribute, it is hypothesized that the reactions occurring within a water-induced, confined organic cage result in a high-pressure-like effect from the surrounding bulk water³⁰. In addition, an enriched hydrogen bond donated by water at the interface is a key factor in facilitating this enhancement³¹. In particular, we discovered that the salting-out effect induced by high-concentration NaCl solutions suppressed micelle formation and increased the interfacial area, further enhancing the reaction efficiency. We hypothesized that this effect could provide a basis for the

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Supplementary Information available: [details of any supplementary information available should be included here]. See DOI: 10.1039/x0xx00000x

efficient functioning of MCFAs as Brønsted acid organocatalysts, particularly below their CMC (Fig. 1B).

Herein, we report the "on-water" accelerated Brønsted acidcatalyzed aqueous reactions by MCFA. As a representative model reaction, the allylboration of a general class of carbonyl compounds was demonstrated, exemplifying the scalability of the gram-scale synthesis. A crucial discovery in this study was the specific salting-out effect induced by NaCl, which proved to be essential for the success of the reaction. Remarkably, the reaction was also effectively catalyzed using very low-cost and unrefined additives such as seawater or edible sea salt, enabling the synthesis of high-purity pharmaceutical intermediates (APIs). The concentrated NaCl solution suppressed the selfassembly of MCFAs as surfactants, thereby enhancing the efficiency of the reaction under "on-water" conditions, specifically at the water-oil interface. This finding highlights the pivotal role of the salting-out effect in facilitating interfacial catalysis and offers a simple and sustainable approach for scalable and cost-effective organic transformations (Fig. 1C).

Result and discussion

Condition Optimization of Catalyst-free Allylation of Aldehydes

We aimed to develop sustainable allylation reactions of general carbonyl compounds (Table 1). For our model reaction, we selected hydrocinnamaldehyde to initiate investigation with aliphatic aldehydes. Allylboronic acid pinacol ester was selected as the allylating reagent owing to its low toxicity and availability³². We first explored conventional organic solvents, such as dichloromethane and acetonitrile stirred at 25 °C. However, only very low conversions to product 1 were obtained (23% and 27% for entries 1 and 2, respectively). The neat condition (no solvent) gave a similarly low reactivity (26%, entry 3). Based on our previous findings, we speculate that aqueous conditions with bulk water as the reaction medium may improve the outcome. Particularly, allylboronic acid pinacol ester is well known to exhibit reasonable activity in aqueous reaction systems^{33,34}. To our surprise, the "on-water" (water-oil biphasic) condition was highly effective in this reaction. The use of pure water (10 L/mol, distilled) significantly improved the results, with >99% conversion in 10 min (entry 6).

Typical slowdown effects were observed under on-water conditions. Utilizing a designer surfactant developed by the Lipshutz group (TPGS-750-M, 2 wt% aqueous solution)³⁵, the entire system was emulsified, yielding a conversion of 87% within 10 min (entry 4). Although this outcome was slightly inferior to that achieved with pure water, the result aligned well with established on-water experimental observations²⁶.

To investigate further, D_2O —a heavier and more viscous medium compared to H_2O —was employed. Similar to the results observed for H_2O , complete conversion was achieved within 10 min. However, the initial reaction rate was relatively slower (86% conversion in 1 min, entry 5) compared to the remarkable 93% conversion achieved within 1 min using H_2O . This notable difference can be attributed to the higher viscosity of D₂O, which likely reduces the effective interfacial surface area between the reactants and the Dadueous³⁹/medium⁴⁴by altering shear forces³¹. The reaction likely proceeds through acceleration at the water–oil interface, in line with the "on-water" theory introduced by Sharpless et al. and later supported theoretically by Jung and Marcus^{26,31}.

Next, NaCl was used as a representative salting-out agent³⁶. This approach yielded exceptional results, achieving >99% conversion within 1 minute (entry 7). These results align with the trends observed in other reported on-water reactions, highlighting accelerated reaction rates due to the effects of aqueous media²⁶. Based on previous findings that hydrophobic agents enhance the efficiency of addition reactions, we replaced the reagent-grade NaCl (from a chemical supplier) with commercially available edible sea salt (from a grocery market). As a result, 95% conversion was achieved within 1 min, and 99% conversion was observed within 2 min (entry 8). Meanwhile, while using bulk water represents an environmentally friendly approach to carbonyl allylation, achieving the initial goal of sustainable conditions and the reliance on distilled water presents challenges. The need for repeated distillation in industrial-scale processes leads to an increased consumption of energy and resources³⁷. From this perspective, seawater, which is one of the most abundant resources on Earth, captured our attention. The utilization of seawater in chemical processes offers significant advantages in terms of cost, toxicity, and environmental impact³⁸. Untreated seawater (salinity = 3.5%) sourced directly from Namhae (the southern sea of Korea, Pohang City) was used as the reaction medium (entry 9). The initial reactivity showed a slightly reduced conversion (84% in 1 min) compared to that achieved with distilled water. However, after 2 min, > 99% conversion was achieved. We believed that the reactivity observed in seawater could be further enhanced by the addition of edible salt. Remarkably, the modification of this medium to saturated seawater (salinity = 26%) significantly improved the reaction rate, achieving 96% conversion within 1 min and complete conversion (>99% conversion) within 2 min (entry 10). These results highlight the potential for utilizing abundant and sustainable resources to develop sustainable and cost-effective synthetic processes.

Based on the optimized reaction conditions, a preliminary investigation of the substrate scope of the aldehydes was conducted (Fig. 2). The reactions were carried out using seawater saturated with edible sea salt as the reaction medium, forming a water-oil biphasic phase. To ensure efficient mixing under these conditions, vigorous stirring (>1000 rpm) was performed²⁹. Owing to the excellent reactivity of the system, all the reactions were completed within 5 min. The model substrate, hydrocinnamaldehyde, afforded product 1 in 99% isolated yield. To further investigate the reactivity of alkyl aldehydes, reactions were carried out with phenylacetaldehyde and 1-cyclohexenylaldehyde, affording products 2 and 3 in high yields of 95% and 90%, respectively. Cinnamaldehyde, a natural product readily obtained from cinnamon bark, afforded product 4 in 97% yield. Furthermore, the reactions with an aryl aldehyde proceeded smoothly, delivering product 5 in exceptionally high yield (99%). To further expand the substrate scope of aryl

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aldehydes incorporating electron-withdrawing and donating substituents, and varying steric effects, were investigated in our reaction system. Para-halogenated substrates (Cl and Br) afforded the corresponding products **6** and **7** in excellent yields (99%). Electron-withdrawing and donating substituents also provided desired products **8–10** in excellent yields (93%–99%). Biphenyl and 2-naphthyl incorporated bulky substrates afforded the desired products **11** and **12** in excellent yields (99%). Moreover, oxygen and sulphur-containing heteroaryl-aldehydes were highly active in the reaction, affording products **13** and **14** in 91% and 99% yields, respectively.

Condition Optimization of Catalytic Allylation of Ketone

We then continued to develop an efficient process for the allylation of less reactive ketones as acceptors³⁹ (Table 2). In particular, tetrasubstituted carbon centers in homoallylic alcohols have attracted our attention because of their potential as key intermediates in the synthesis of various active APIs^{40,41}. Conventional methods such as the Hosomi–Sakurai allylation employing organosilicon reagents such as allyl trimethylsilane in moisture-sensitive metal Lewis acids have been widely investigated^{42,43}. However, the application of anhydrous conditions to industrial processes poses significant challenges. The avoidance of organic solvents and metals in this catalytic system was expected to have a substantial impact.

Acetophenone, an aryl-alkyl ketone, was selected as the model substrate, and water was used as the reaction medium. Unlike aldehydes, ketones exhibited significantly lower reactivities under catalyst-free conditions, with negligible formation of product **15** (entry 1). At an elevated temperature of 60 °C, the reaction proceeded with 29% conversion after 24 h of stirring (entry 2). Consequently, we hypothesized that either Brønsted or Lewis acid catalysts would be required and proceeded to screen various candidates.

Inspired by our previous work on multicomponent aquacatalytic allylation, we preferably evaluated strong Brønsted acids²¹. *p*-toluenesulfonic acid However, (PTSA), dodecylbenzenesulfonic acid (DBSA), and bistriflimide led to the complete decomposition of the substrate (~0%), whereas triflic acid (TfOH) showed limited catalytic activity (38% conversion). Among Lewis acids, tris(pentafluorophenyl)borane (B(C₆F₅)₃), a highly effective catalyst for Mukaiyama aldol reactions in water reported by the Loh group⁴⁴, was inactive in this system. Scandium triflate (Sc(OTf) 3), commonly utilized by the Kobayashi group⁴⁵, demonstrated marginally better activity but suboptimal performance, with a conversion of 54%. These findings highlight the need for novel and efficient catalysts.

Carboxylic acids are notable for their natural abundance, straightforward production, mild acidity, and low toxicity, all of which contribute to their safety and economic viability⁴⁶. A diverse range of carboxylic acids was screened as catalysts (10 mol%). Simple highly water-soluble acids, such as formic acid (Log P = -0.54) and acetic acid (Log P = -0.26), exhibited no catalytic effect (31–33% conversion), while trifluoroacetic acid (Log P = 1.35), benzoic acid (Log P = 1.95), and *p*-dodecylbenzoic acid showed almost no reactivity (<1% conversion, entries 3–7). Considering the aqueous reaction medium, we hypothesized

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that introducing longer alkyl chains into the carboxylic acid structure might enhance the reactivity O^{1} Natural Dfatty 14cid screening revealed promising results. Caproic acid (C6:0, Log P = 1.92), caprylic acid (C8:0, Log P = 3.05), capric acid (C10:0, Log P = 4.09) and lauric acid (C8:0, Log P = 5.15) which are key components of MCTs³, significantly improved the conversion to 40%, 53%, 48% and 45% (entries 8–11). In contrast, longer-chain fatty acids such as myristic acid (C14:0, Log P = 6.10) and erucic acid (C22:1, Log P = 9.46), as well as branched acids such as isobutyric acid (Log P = 0.45), showed no further improvement (entries 12–14).

Among the tested acids, caprylic acid emerged as the optimal catalyst owing to its lipophilicity (Log P = 3.05). Temperature also played a critical role in the reaction. At 80 °C, the reactivity increased moderately, achieving 66% conversion (entry 15), while at 100 °C, the conversion decreased. This decrease was probably due to the increased solubility of the fatty acid above its CMC¹⁶, which disrupted the biphasic reaction environment for catalysis (entry 16). The salting-out effect of NaCl is particularly crucial in this system, as it further enhances the reactivity. Under the optimized conditions, increasing the catalyst loading to 30 mol% led to the complete conversion to the desired product **15** (entries 17–20, and 22).The reaction yield dropped dramatically to 25% when sodium caprylate was used (entry 20).

Substrate Scope of Allylation of Ketone

Based on the optimized reaction conditions, the substrate scope for the allylation of various ketones was explored (Fig. 3). Ketones substituted with alkyl-alkyl groups are generally less reactive as acceptors, presenting a significant challenge. However, under our salting-out-enhanced "on-water" conditions, they were successfully converted to compounds 16 and 17 in excellent yields (97% and 91%, respectively). The method was further extended to cyclic ketones, yielding products 18 and 19 with remarkable efficiencies (up to 95% yield). Similarly, alkenyl and alkynyl ketones underwent quantitative transformations into the expected products, 20 and **21**, in 99% and 96% yields, respectively. Substrates bearing (thio)ester functional groups, such as β -ketoesters and α ketothioesters, also exhibited excellent functional group tolerance, yielding compounds 23 and 24 in 97% and 90% yields, respectively. A wide range of (hetero)aryl-alkyl ketones were also converted to the desired products without complications in very high yields (compounds 15-38, up to 99%). Additionally, gram-scale reactions with ketone substrates were successfully conducted, maintaining excellent efficiency (99% yield for compounds 15 and 34).

We further demonstrated the utility of this method by performing late-stage modifications on ketone-containing natural products and pharmaceuticals. Nabumetone⁴⁷, a nonsteroidal anti-inflammatory drug (NSAID), and flavone⁴⁸, a compound commonly found in plant-derived foods, were efficiently transformed into the desired products in high yields (99% yield for compound **39**; 98% yield with anti/syn = 3.7/1 for compound **40**). This investigation highlights the broad applicability and robustness of the developed aqueous

allylation protocol, even with challenging substrates and latestage functionalization.

Development of Chromatography-Free Purification Methods and Synthetic Applications

To achieve an efficient and sustainable purification process, it is necessary to develop a purification protocol that does not require column chromatography. A 1.0-gram scale synthesis of benzylacetone was conducted to isolate the product solely through extraction. The reaction was performed using untreated seawater saturated with edible sea salt as a medium (Fig. 4A). Upon completion of the reaction, the crude mixture was subjected to extraction-based purification. The most challenging aspect of the purification process was the removal of the residual allyl-Bpin and caprylic acid. Previous studies have indicated that pinacol boronate esters undergo hydrolysis under acidic conditions⁴⁹. Subsequently, 6 N HCl (aq.) was added to the crude mixture, which was then stirred for 24 h. This treatment transformed the pale-yellow crude mixture into a dark red solution, and the resulting unreacted components were simply removed through extraction. Next, the removal of residual caprylic acid, which was used as the catalyst, was investigated. The water solubility of caprylic acid increases with temperature⁵⁰, allowing its efficient removal through hot water extraction (see Supporting Information). To validate the efficacy of this protocol, gas chromatography (GC) was performed on both crude and purified products. The residual allyl-Bpin, caprylic acid, and minor benzylacetone impurities were removed entirely, yielding a highly pure (>99%) GC trace of product 16. This successful chromatography-free protocol is also enabled to the aldehyde (see Supporting Information) and highlights the potential for simplifying sustainable and economical processes.

To further demonstrate the utility of these reaction products, olefin metathesis was performed using **15** (Fig. 4B). Using the Grubbs-II catalyst, compound **15** was successfully converted to compound **41** with 80% yield. This excellent functional group tolerance underscores the potential of this method for API synthesis.

The compatibility of our catalytic allylation system with water and its transition-metal-free nature suggest that it is particularly practical for biomolecule functionalization applications⁵¹. In this context, the reaction system aligns closely with the biomimetic conditions required for constructing DNA-encoded libraries (DELs), a powerful tool in drug discovery⁵². Notably, the allylation of ketones tethered to DNA has not yet been reported, making this a significant opportunity for advancing its biochemical applications. To extend the applicability of this reaction, we investigated its use in on-DNA allylation (Fig. 4C). First, amide coupling was performed by reacting a headpiecemodified DNA molecule⁵³ with 4-acetylbenzoic acid, hexafluorophosphate azabenzotriazole tetramethyl uranium (HATU), N,N-diisopropylethylamine (DIPEA), and pH 9.5 borate buffer at 80 °C for 24 h. Compound 42 was characterized using HPLC and UPLC, achieving a conversion of 92% (HRMS: m/z calcd. $[M - H]^- = 5327.0540$; found $[M - H]^- = 5327.1348$). Compound 42 was diluted with double-distilled H2O. Allyl-Bpin

(500 equiv.) and caprylic acid (100 equiv.) were added, and the reaction was performed at 80 °C for 24 R. The desired product **43** was successfully obtained with 56% conversion, as confirmed by HPLC and UPLC (MS: m/z calcd. $[M - H]^- = 5369.1010$; found $[M - H]^- = 5369.1750$). This study highlights the utility of the developed catalytic system for sustainable and efficient aquatic allylation processes and its broad applicability to advanced biochemical transformations such as DNA-encoded library synthesis

Mechanistic investigation

Investigating the water-promoted rate acceleration in biphasic reactions involving a catalyst is inherently more complex than in catalyst-free conditions. The mechanism underlying this enhancement was well categorized by Kobayashi and Kitanosono^{54,55}, who classified on-water reactions involving catalysts as type III. In our system, elevated temperatures increase the solubility of caprylic acid in water; however, the salting-out effect induced by saturated Na⁺ and Cl⁻ ions may be expected to suppress micelle formation (illustrated in Fig. 1C). It is hypothesized that the reaction predominantly occurs at the interface between the oil phase (the ketone, allyl boronate, and caprylic acid (Log P = 3.05)) and the ion-saturated aqueous phase. Jung and Marcus proposed that the free –OH groups in bulk water form stronger hydrogen bonds with hydrogen-bond acceptors in the transition state³¹. Additionally, based on computational studies, Roke et al. reported that the surface of pure water is more acidic than the bulk⁵⁶, suggesting that the acidity of hydrated caprylic acid is enhanced at the oil-water interface. This increased acidity may facilitate strong hydrogen bonding interactions with hydrogen bonding acceptors such as the oxygen atom adjacent to the boron atom in allyl-Bpin^{57,58}. To further elucidate this phenomenon, density functional theory (DFT) calculations were performed (Fig. 5A). A key aspect of this investigation was to determine whether the catalytic activity of caprylic acid could be attributed to its -COOH group, which activates allyl-Bpin to provide the [intermediate-I]. DFT calculations revealed that the Mulliken charge of the boron atom in allyl-Bpin increased from 1.645 in the absence of a catalyst to 1.961 in the presence of caprylic acid. This suggests that the caprylic acid forms a hydrogen-bonded complex with allyl-Bpin, enhancing its Lewis acidity and generating additional reactive organoboron intermediates. For comparison, further calculations were performed to examine in the case of the interaction between caprylic acid and the carbonyl oxygen of acetophenone ([intermediate-II]). As a result, the Mulliken charge on the carbonyl carbon of acetophenone increased from 0.126 to 0.246 upon interaction with caprylic acid. This outcome suggests that caprylic acid preferentially interacts with the oxygen atom in allyl-Bpin rather than with the carbonyl oxygen in acetophenone (see Supporting Information for detail).58

To validate whether the reactive intermediate benefited from the on-water effect, the reactivities of various reaction media were compared (Fig. 5B). Reactions were conducted with 10 mol% caprylic acid at an optimized temperature of 80 °C. Bulk water yielded 66% conversion (entry h), whereas organic solvents such as acetone and dichloromethane were completely

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inactive. Methanol exhibited a limited reactivity (16% conversion; entries a-c). Although the initial rate was slightly slower, D_2O gave a result comparable to that of H_2O . (entry g). The designer surfactant H₂O/TPGS (2 wt%) resulted in only 30% yield (entry d), while neat conditions improved the yield to 62% (entry f). These findings indicate that homogeneous reaction systems are unsuitable, with their reaction performance closely linked to substrate concentration; higher concentrations under neat conditions enhance the reaction. The superior reactivity observed in bulk water is likely derived from the biphasic system, which creates a confined organic cage that exerts a highpressure-like effect on the reactants³⁰. This confinement compresses the reactive organic phase, leading to a negative activation volume (ΔV^{\dagger} < 0), which is known to accelerate transformations, particularly those involving addition-type reaction mechanisms²⁹. This hypothesis is consistent with the observed trends: a hydrophobic agent³⁴ like NaCl enhances conversion to 79% (entry e), while an anti-hydrophobic agent⁵⁹ like LiClO₄ reduces it to 52% (entry d). Furthermore, increasing the catalyst loading significantly improves the reactivity, highlighting the critical role of the catalyst (entries j-m).

To ensure that the catalytic system followed a Zimmerman-Traxler-type transition state⁶⁰, we replaced allyl-Bpin with *trans*-crotylboronic acid pinacol ester (compound **44**, Fig. 5C). The reaction yielded only compound **45**, with a methyl group at the α -position in 95% yield. Notably, a single diastereomer with an *anti*-configuration was observed, indicating that the nucleophilic attack on the carbonyl carbon occurred exclusively at the terminal sp²-carbon atom. In accordance with our expectations, reactions conducted in organic solvents (MeCN) afforded less than 5% yield, and even under solvent-free conditions, the yield was low (60%). These results support the involvement of the Zimmerman–Traxler transition state.

Conclusions

In summary, we have developed a sustainable aqueous catalytic approach for carbonyl allylation, leveraging unrefined seawater and sea salt to harness the salting-out effect. This method achieved (i) efficient, catalyst-free allylation of aldehydes and (ii) caprylic acid-catalyzed allylation of ketones using a costeffective, nontoxic fatty acid commonly employed in the food industry, with bulk water as the reaction medium. The process demonstrated scalability, enabling gram-scale synthesis with chromatography-free purification and offering a promising pathway for cost-efficient API production. Additionally, the mild reaction conditions and water compatibility facilitate on-DNA conjugation, potentially transforming the development of DELs, a critical innovation in drug discovery. We believe that this study provides a robust foundation for further advancements in environmentally friendly, water-compatible, and fundamental catalytic systems for challenging applications.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

View Article Online DOI: 10.1039/D5GC01441G

The generous support of the Ministry of Science, ICT, and Future Planning of Korea (RS-2023-00259659 and RS-2023-00219859) and the Ministry of Education (2022R1A6C101A751 and 2022R1A6C102A913) is gratefully acknowledged. H.Y.B. thanks Korea Toray Fellowship.

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Fig. 1 Concept underlying this study. A Structure and properties of MCFA. B Activated Brønsted acid catalysis under concentrated salty aqueous medium. C "On-Seawater" accelerated aquacatalysis harnessing the remarkable salting-out effect.

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Table 1 Medium effect on the catalyst-free allylboration of aldehyde^a

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Entry	[reaction medium]	Conv. (%) ^b				
		1 min	2 min	3 min	5 min	10 min
1	CH ₂ Cl ₂	3	5	8	18	23
2	MeCN	2	3	10	20	27
3	neat	3	5	9	20	26
4	TPGS-750-M (2 wt%)	73	75	79	83	87
5	D ₂ O	86	>99	>99	>99	>99
6	H ₂ O	93	>99	>99	>99	>99
7	H ₂ O-NaCl (sat.)	>99	>99	>99	>99	>99
8	H ₂ O-edible salt (sat.)	95	>99	>99	>99	>99
9	Seawater	84	>99	>99	>99	>99
10	Seawater-edible salt (sat.)	96	>99	>99	>99	>99

^{*a*}Reactions were conducted using hydrocinnamaldehyde (0.1 mmol, 1.0 equiv.) and allyl-Bpin (0.12 mmol, 1.2 equiv.) in different reaction media (10 L/mol) at 25 °C. ^{*b*}Conv. (%) was determined by ¹H NMR using 1,4-dimethoxybenzene as an internal standard. min = minute.

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View Article Online DOI: 10.1039/D5GC01441G catalyst-free HO seawater-edible salt (sat.) vigorous stirring (>1000 rpm) 25 °C [aldehydes] 5 min 1-14 ΗQ НQ HO н HO H н н но **3** 90% yield **2** 95% yield **5** 99% yield **4** 97% yield **1** 99% yield HO н HO н H HO н HO н HO CI Br F₃C O₂N BnO 6 8 9 10 7 99% yield 99% yield 93% yield 99% yield 96% yield HO H но н HQ Н но н Ph 11 **12** 99% yield **13** 91% yield **14** 99% yield 99% yield Mg²⁺Ca²⁺ K⁺ seawater-edible salt (sat.) edible salt seawater SO4 청정원 CI-순수천혜염 Na 55.1 일염가는소금 30.6 six major ions in sea salt (wt%)

Fig. 2 Substrate diversity of the catalyst-free allylboration of aldehydes. Reactions were conducted using aldehyde (0.2 mmol, 1.0 equiv.), allyl-Bpin (0.24 mmol, 1.2 equiv.) and seawater-edible salt (sat.) at 25 °C for 5 min.

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acid catalyst (10-30 mol%)

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	$\sum_{F} B(C_6F_5)_3$	triflic acid	Sc(OTf) ₃		
	· F decomposed	conv. = 38%	conv 54 %		
Entry	carboxylic acid or salt (mol%)	Log P of acid	[reaction medium]	Conv. (%) of 5 ^b	
1 ^{<i>c</i>}	without catalyst	-	H ₂ O	5	
2	without catalyst	-	H ₂ O	29	
3	formic acid (10)	-0.54 (-0.30)	H ₂ O	33	
4	acetic acid (10)	-0.26 (-0.09)	H ₂ O	31	
5	trifluoroacetic acid (10)	1.35 (0.21)	H ₂ O	<1	
6	benzoic acid (10)	1.95 (1.65)	H ₂ O	<1	
7	4-n-dodecylbenzoic acid (10)	(6.81)	H ₂ O	<1	
8	caproic acid (10)	1.92 (1.83)	H ₂ O	40	
9	caprylic acid (10)	3.05 (2.93)	H ₂ O	53	
10	capric acid (10)	4.09 (4.04)	H₂O	48	
11	lauric acid (10)	5.15 (4.60)	H ₂ O	45	
12	isobutyric acid (10)	0.94 (0.45)	H ₂ O	43	
13	myristic acid (10)	6.10 (4.97)	H ₂ O	32	
14	erucic acid (10)	9.46 (8.10)	H ₂ O	10	
15 ^d	caprylic acid (10)	3.05 (2.93)	H ₂ O	66	
16 ^e	caprylic acid (10)	3.05 (2.93)	H ₂ O	trace	
17 ^d	caprylic acid (10)	3.05 (2.93)	H ₂ O-NaCl (sat.)	79	
18 ^d	caprylic acid (20)	3.05 (2.93)	H ₂ O-NaCl (sat.)	87	
19 ^a	caprylic acid (30)	3.05 (2.93)	H₂O-NaCl (sat.)	92	
20 ⁷	caprylic acid (30)	3.05 (2.93)	H ₂ O	83	
21 ⁷	sodium caprylate (30)	-	H ₂ O-NaCl (sat.)	25	
22	caprylic acid (30)	3.05 (2.93)	H ₂ O-NaCl (sat.)	>99(99) ^g	

^oReactions conducted on 7 (0.1 mmol, 1.0 equiv.), allyl-Bpin (0.3 mmol, 3.0 equiv.) and a carboxylic acid catalyst (10 mol%) in H₂O (10 L/mol) at 60 °C for 24 h. ^bConv (%) was determined by ¹H NMR analysis using 1,4-dimethoxybenzene as an internal standard. ^cAt 25 °C. ^dAt 80 °C. ^eAt 100 °C. ^fUsing allyl-Bpin (5.0 equiv.) at 80 °C. ^gIsolated yields are given in parentheses.

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Fig. 3 Substrate scope of the fatty acid-catalyzed allylboration of ketones. Reactions were conducted using ketone (0.2 mmol, 1.0 equiv.), allyl-Bpin (1.0 mmol, 5.0 equiv.), caprylic acid (30 mol%), and H_2O -NaCl (sat., 2.0 mL) at 80 °C for 24 h. ^{*a*}48 h of reaction time. ^{*b*}Isolated yield of gram-scale synthesis.

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Fig. 4 Synthetic utilities. **A** Reaction conducted using benzylacetone (1.0 g), allyl-Bpin(3.0 equiv.), caprylic acid (30 mol%), and seawateredible salt (sat.) at 80 °C for 24 h. Description of pictures: (i) Concentrated crude mixture after reaction completion. (ii) Addition of 6 N HCl (aq.). (iii) Extraction with ethyl acetate/water to remove organic impurity. (iv) Extraction with ethyl acetate/hot water to remove the remaining caprylic acid catalyst. (v) Obtained desired product **16** (>99% GC purity). **B** Derivatization: catalytic olefin metathesis of the product. **C** On-DNA bioconjugation process.



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Fig. 5 Mechanistic investigation. **A**. Proposed catalytic cycle and DFT-calculated Mulliken charges of selected atoms. **B** Effect of reaction medium on catalytic allylboration of acetophenone. Reactions were conducted using acetophenone (0.1 mmol, 1.0 equiv.), allyl-Bpin (0.3 mmol, 3.0 equiv.), caprylic acid (10 or 30 mol%), and the reaction medium at 80 °C for 24 h. Conv (%) were determined by ¹H NMR analysis using 1,4-dimethoxybenzene as an internal standard. **C** Validation of Zimmerman-Traxler mediated mechanism. The reaction was conducted using acetophenone (0.1 mmol, 1.0 equiv.), compound **44** (0.5 mmol, 5.0 equiv.), caprylic acid (30 mol%), and H₂O-NaCl (sat.) at 80 °C for 24 h.

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Data availability

The data supporting the findings of this study are available within the article and its Supplementary Information. Additional data are available from the corresponding author upon request.