



Late-Stage C–H Acylation of Tyrosine-Containing Oligopeptides with Alcohols

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technique is distinguished by its reliable scalability and features the use of ethanol as a renewable feedstock for the assembly of a variety of peptidomimetics.

S ince the definition of the "12 Principles of Green Chemistry" by Anastas and Warner in 1998,¹ sustainable development represents a major global concern when designing new chemical synthetic processes.² As a result, the last decades have witnessed the upsurge of a sheer number of greener and safer procedures for the synthesis of fine and commodity chemicals. In particular, the practical use of abundant and renewable carbon feedstock in the realm of organic chemistry has gained considerable attention.³ However, the use of ethanol as a valuable and cheap C₂ feedstock is still rare, and it is chiefly used as an organic solvent rather than an actual coupling partner.⁴ In this communication, we unlock its synthetic versatility and advantageous features within the burgeoning field of bioconjugation.

Owing to their unique biological activities and improved metabolic stability compared to their native compounds, synthetically modified peptides are of utmost importance in the field of proteomics, chemical biology, and drug discovery.⁵ Metal catalysis has recently emerged as an enabling tool for the manipulation of typically unreactive C-H bonds embedded within the amino acid backbone⁶ and the corresponding side chains.' Accordingly, metal-catalyzed C-H functionalization techniques are becoming highly embraced by mainstream synthetic chemists because they enable the straightforward assembly of biomolecules in a sustainable fashion.⁸ Despite the existing palette of reactivity, most of the protocols entail the use of toxic halide counterparts and feature the modification of highly reactive amino acid residues. Therefore, innovative tactics are highly coveted to forge peptides beyond those found in naturally occurring proteins, and the usage of new atomeconomical C-H coupling partners to label less reactive and poorly nucleophilic handles represents an ideal strategy in these endeavors. The modification of peptides housing hydrophobic phenylalanine (Phe) and tyrosine (Tyr) residues remains comparatively overlooked,⁹ which is clear evidence that the direct translation of a given $C(sp^2)$ -H functionalization reaction from a simple aryl system to a peptide framework is not a trivial task as a result of the existing multiple chelating sites and ubiquitous C–H bonds. 10

Recent studies have demonstrated that the installation of an acetyl group within an amino acid of a peptide sequence is particularly useful to produce antibody-drug conjugates through oxime ligation.¹¹ Although acetylated proteins are primarily prepared upon enzymatic processes with acetyl-transferases or acetyl-CoA derivatives,¹² the parent processes in short-to-medium peptides remain elusive. The orthoacetylation of simple L-Tyr-OH can occur through a classical Friedel-Crafts reaction with acetyl chloride.¹³ However, the latter cannot be applied within a peptide setting. Partial racemization is often observed (up to 15%), and stoichiometric amounts of AlCl₃ are required (Scheme 1). In connection with our previous studies on the modification of peptides,¹⁴ we sought to tackle the synthetic potential of EtOH as a novel acetyl source under oxidative conditions, thereby providing a sustainable yet late-stage acetylation of a number of Tyrcontaining compounds. While conceptually innovative, this strategy may suffer from certain drawbacks, such as the lack of selectivity or even an undesired ortho-alkoxylation reaction could preferentially occur when using EtOH.¹⁵ Herein, we present a complementary strategy to perform a chloride-free acetylation of Tyr-containing peptides, which can take place in a late-stage fashion featuring cheap and safe chemical reagents.

Inspired by the use of 2-pyridyl ether as an efficient directing group $(DG)^{16,17}$ in the Pd-catalyzed acylation of protected Tyr derivatives with aldehydes recently reported by our group,^{10c} we first selected dipeptide 1a as the model substrate to test the

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Scheme 1. ortho-Acetylation of Tyr Derivatives







Table 1. Pd-Catalyzed C–H Acetylation of Compound 1a with Ethanol^a



^{*a*}Reaction conditions: compound 1a (0.15 mmol), EtOH (3.75 mmol, 0.2 mL), $Pd(OAc)_2$ (10 mol %), and T-hydro (6.0 equiv) in PhMe (1 mL) at 120 °C for 16 h under Ar. T-hydro = *tert*-butyl hydroperoxide solution, 70 wt % in water; DCP = dicumyl peroxide. ^{*b*}Yield of isolated product after column chromatography.

feasibility of the acetylation reaction with EtOH, thereby harnessing its oxidation toward the sustainable labeling of peptides. After considerable experimentation,¹⁸ we eventually found that the projected acetylation with EtOH was feasible, and remarkably, neither diacetylation nor ortho-alkoxylation upon a C–O bond-forming event was detected.¹⁵ The optimal conditions involved the use of $Pd(OAc)_2$ (10 mol %) and an aqueous solution of inexpensive tert-butyl hydroperoxide (TBHP) as the oxidant in toluene as the solvent at 120 °C, which provided compound 2aa in 60% yield (entry 1 of Table 1). Notably, toluene was not activated to produce the corresponding aroylated product, and EtOH was preferentially oxidized within the reaction conditions.¹⁹ As expected, control experiments in the absence of either catalyst (entry 2 of Table 1) or oxidant (entry 3 of Table 1) underpinned their critical role in the acetylation process. The performance of the reaction under air resulted in lower yields of compound 2aa,

Scheme 2. Pd-Catalyzed C–H Acylation of Compound 1a with Alcohols a,b



^{*a*}Reaction conditions: compound **1a** (0.15 mmol), RCH₂OH (0.75 mmol), Pd(OAc)₂ (10 mol %), and T-hydro (6.0 equiv) in PhMe (1 mL) at 120 °C for 16 h under Ar. ^{*b*}Yield of isolated product after column chromatography, with the average of at least two independent runs. ^{*c*}Reaction conditions: compound **1a** (0.15 mmol), RCH₂OH (0.45 mmol), Pd(OAc)₂ (10 mol %), and T-hydro (4.0 equiv) in PhMe (1 mL) at 120 °C for 16 h under Ar.

albeit the process still occurred in a synthetically relevant yield (entry 4 of Table 1). Whereas the yield slightly dropped down to 40% when using 10 equiv of EtOH (entry 5 of Table 1), the process was entirely inhibited in EtOH as the solvent (entry 6 of Table 1). Accordingly, the optimal amount of EtOH was found to be 25 equiv in combination with toluene as the solvent; the use of other related solvents ushered compound 2aa in lower yields.¹⁸ Given that multiple oxidation events simultaneously occur, the yield reasonably decreased when lowering the amount of TBHP (entries 7 and 8 of Table 1). However, its use in high excess does not pose a major shortcoming because it is a very cheap oxidant and renders the reaction water-compatible. In fact, an aqueous solution of TBHP afforded better results than other peroxides or persulfates (entries 9 and 10 of Table 1), and $Pd(OAc)_2$ clearly outperformed other palladium catalysts¹⁸ (entries 11 and 12 of Table 1). Finally, we confirmed that subtle modifications on the DG had a determinant impact on the reaction outcome, and the OPy motif was the most active DG toward the target acetylation reaction (Table S3 of the Supporting Information).^{16,18}

Although we primarily focused on the unprecedented use of EtOH to acetylate peptides in a site-selective manner, we also evaluated the parent acylation process of dipeptide 1a using other related aliphatic alcohols. For instance, inexpensive *n*-BuOH, 4-methyl-1-pentanol, and even biologically relevant palmityl alcohol derived from the corresponding fatty acid



Scheme 3. Pd-Catalyzed C-H Acylation of Tyr-Containing Oligopeptides with EtOH and Other Alcohols^{*a,b*}

^{*a*}The same as for entry 1 of Table 1. ^{*b*}Yield of isolated product after column chromatography, with the average of at least two independent runs with a variable yield by no more than 5% between runs. ^{*c*}Using PhCl instead of PhMe as the solvent. ^{*d*}Compound 1 (0.15 mmol), alcohol (0.45 mmol), Pd(OAc)₂ (10 mol %), and T-hydro (4.0 equiv) in PhMe (1 mL) at 120 °C for 16 h under Ar. ^{*e*}Compound 1 (0.15 mmol), alcohol (0.75 mmol) Pd(OAc)₂ (10 mol %), and T-hydro (6.0 equiv) in PhMe (1 mL) at 120 °C for 16 h under Ar.

resulted in the exclusive monoacylated products **2ab-2ad** in high yields (Scheme 2).

Noteworthy, in those cases, the amount of alcohol could be significantly reduced to 5.0 equiv. Likewise, activated benzyl alcohols^{20,21} could also be employed to produce the corresponding aroylated dipeptides in 59-63% yields. Owing to their higher tendency toward oxidation, the reaction conditions were slightly modified to avoid the formation of the diaroylated compound; thus, lower amounts of both oxidant and alcohol were required. These experiments revealed that benzyl alcohols could be practical surrogates of

benzaldehydes to perform the *ortho*-acylation of Tyr compounds,^{10c} thereby providing exclusively the monofunctionalized products.

We next explored the synthetic scope of the acetylation manifold featuring EtOH in the challenging setting of short-tomedium-size peptides (Scheme 3). Notably, peptides bearing Val (1c), Phe (1d), Lys (1e), Ala (1f), Pro (1g), Gly (1h), Ser (1i), Asp (1j), Glu (1k), Ile (1l), Tyr (1m), and even Arg (1p) were found compatible with the reaction conditions and provided the corresponding acetylated peptides in moderate to good yields. Note that the N terminus and other oxidizable

Scheme 4. Gram-Scale Synthesis



amino acid residues housing a free amino, an alcohol, a carboxylic acid, or a guanidine motif (Lys, Ser, Asp, Glu, and Arg, respectively) were equipped with protecting groups to achieve chemoselectivity. Notably, this labeling technique was applicable to Tyr residues located at the N and C terminals as well as inner positions. Importantly, tetrapeptide 1n and hexapeptide 10 having the sequence of biologically relevant endomorphin-2 and neuromedin N, respectively, were also acetylated with EtOH, hence showcasing the high utility of this method toward the site-selective tagging of complex biomolecules. As previously anticipated, other alcohols could also be selectively installed at the ortho position of the Tyr unit within di-, tri-, and tetrapeptide derivatives (2db, 2kb, and 2q and 2r). In general, the reactions were very clean, and side products were not observed, albeit full conversion was not always achieved and sometimes PhMe was replaced by more oxidizing PhCl. Besides, unlike classical Friedel-Crafts acetylation, our method features the use of EtOH as a sustainable C₂ source to accomplish a synthetically meaningful transformation, wherein a high number of C-H bonds are activated. In this respect, the acylation of compound 1a could be performed in gram scale when using EtOH and BuOH with a remarkable 62 and 74% yield, respectively (Scheme 4). In these cases, the amount of EtOH and oxidant could be slightly reduced without affecting the reaction outcome, which represents a promising starting point for applied research.

To gain some insights into the reaction mechanism, we conducted some control experiments. We found that the acetylation of compound 1a was suppressed in the presence of 2,2,6,6-tetramethylpiperidine 1-oxyl (TEMPO), which indicated that a radical pathway may be operative.¹⁸ Furthermore, assuming that EtOH could be oxidized to acetaldehyde within the course of the reaction, we performed some tests with MeCHO as the coupling partner.¹⁸ When using acetaldehyde under the acylation conditions previously developed by our group involving water as the solvent, 10c traces of compound 2aa were obtained, which reveals the subtleties of installing a simple acetyl group. Notably, the use of PhMe as the solvent resulted in mixtures of mono- and diacetylated products, and the use of a high excess of MeCHO ushered in the exclusive formation of diacetylated compound 2aa' in 62% yield (Table S4 of the Supporting Information).¹⁸ Accordingly, if EtOH is in situ transformed into MeCHO in the presence of TBHP,²² the reaction mechanism should be akin to those of related acylations with aldehydes described in the literature.^{18,21} The high selectivity toward the monoacetylation could be due to the lower reactivity of EtOH in comparison to the corresponding aldehyde.

In summary, we have demonstrated the high versatility of EtOH as a sustainable feedstock to tag Tyr-containing peptides in a late-stage fashion. This reliably scalable platform represents an innovative avenue for the diversification of Tyr-containing compounds. Salient features of this method are the widespread availability and low cost of EtOH and other related alcohols, the compatibility with an aqueous environment, and the site-selectivity toward the monofunctionalization of the Tyr unit within a peptide setting. Accordingly, this Pd-catalyzed acetylation manifold represents a useful tool for the facile modification of a virtually unlimited number of biologically relevant peptides.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.orglett.1c02764.

Experimental procedures, syntheses and characterization of all new compounds, and tables with details of several optimization studies (PDF)

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Notes

The authors declare no competing financial interest.

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REFERENCES

(1) Anastas, P.; Eghbali, N. Green Chemistry: Principles and Practice. *Chem. Soc. Rev.* 2010, 39, 301.

(2) (a) Schaub, T. Efficient Industrial Organic Synthesis and the Principles of Green Chemistry. *Chem. - Eur. J.* 2021, 27, 1865.
(b) Dalton, T.; Faber, T.; Glorius, F. C-H Activation: Toward Sustainability and Applications. *ACS Cent. Sci.* 2021, 7, 245.

(3) (a) Kühlborn, J.; Groß, J.; Opatz, T. Making Natural Products from Renewable Feedstocks: Back to the Roots? *Nat. Prod. Rep.* **2020**, 37, 380. (b) Tortajada, A.; Juliá-Hernández, F.; Börjesson, M.; Moragas, T.; Martin, R. Transition-Metal-Catalyzed Carboxylation

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Reactions with Carbon Dioxide. Angew. Chem., Int. Ed. 2018, 57, 15948.

(4) (a) Meyer, C. C.; Stafford, N. P.; Cheng, M. J.; Krische, M. J. Ethanol: Unlocking an Abundant Renewable C2-Feedstock for Catalytic Enantioselective C-C Coupling. *Angew. Chem., Int. Ed.* **2021**, 60, 10542. (b) Xie, Y.; Ben-David, Y.; Shimon, L. J. W.; Milstein, D. Highly Efficient Process for Production of Biofuel from Ethanol Catalyzed by Ruthenium Pincer Complexes. *J. Am. Chem. Soc.* **2016**, *138*, 9077. (c) Aitchison, H.; Wingad, R. L.; Wass, D. F. Homogeneous Ethanol to Butanol Catalysis-Guerbet Renewed. ACS Catal. **2016**, *6*, 7125. (d) Wang, G.-W.; Yuan, T.-T. Palladium-Catalyzed Alkoxylation of N-Methoxybenzamides via Direct sp² C-H Bond Activation. J. Org. Chem. **2010**, *75*, 476.

(5) (a) Lenci, E.; Trabocchi, A. Peptidomimetic Toolbox for Drug Discovery. *Chem. Soc. Rev.* **2020**, *49*, 3262. (b) Lau, J. L.; Dunn, M. K. Therapeutic peptides: Historical Perspectives, Current Development Trends, and Future Directions. *Bioorg. Med. Chem.* **2018**, *26*, 2700.

(6) San Segundo, M.; Correa, A. Cross-Dehydrogenative Coupling Reactions for the Functionalization of α -Amino Acid Derivatives and Peptides. *Synthesis* **2018**, *50*, 2853.

(7) (a) King, T. A.; Kandemir, J. M.; Walsh, S. J.; Spring, D. R. Photocatalytic Methods for Amino Acid Modification. Chem. Soc. Rev. 2021, 50, 39. (b) Zhan, B.-B.; Jiang, M.-X.; Shi, B.-F. Late-Stage Functionalization of Peptides via a Palladium-Catalyzed C(sp³)-H Activation Strategy. Chem. Commun. 2020, 56, 13950. (c) Tong, H.-R.; Li, B.; Li, G.; He, G.; Chen, G. Postassembly Modifications of Peptides via Metal-Catalyzed C-H Functionalization. CCS Chem. 2021, 3, 1797. (d) Guerrero, I.; Correa, A. Site-Selective Trifluoromethylation Reactions of Oligopeptides. Asian J. Org. Chem. 2020, 9, 898. (e) Wang, W.; Lorion, M. M.; Shah, J.; Kapdi, A. R.; Ackermann, L. Late-Stage Peptide Diversification by Position-Selective C-H Activation. Angew. Chem., Int. Ed. 2018, 57, 14700. (f) Noisier, A. F. M.; Brimble, M. A. C-H Functionalization in the Synthesis of Amino Acids and Peptides. Chem. Rev. 2014, 114, 8775. (8) (a) Guillemard, L.; Kaplaneris, N.; Ackermann, L.; Johansson, M. J. Late-Stage C-H Functionalization Offers New Opportunities in

Drug Discovery. Nat. Rev. Chem. 2021, 5, 522. (b) Cernak, T.; Dykstra, K. D.; Tyagarajan, S.; Vachal, P.; Krska, S. W. The Medicinal Chemist's Toolbox for Late Stage Functionalization of Drug-Like Molecules. Chem. Soc. Rev. 2016, 45, 546. (c) Gutekunst, W. R.; Baran, P. S. C-H Functionalization Logic in Total Synthesis. Chem. Soc. Rev. 2011, 40, 1976.

(9) (a) Correa, A. Metal-Catalyzed $C(sp^2)$ -H Functionalization Processes of Phenylalanine- and Tyrosine-Containing Peptides. *Eur. J. Inorg. Chem.* **2021**, 2021, 2928. (b) Dorta, D. A.; Deniaud, D.; Mével, M.; Gouin, S. G. Tyrosine Conjugation Methods for Protein Labelling. *Chem.* - *Eur. J.* **2020**, 26, 14257. (c) Szijj, P. A.; Kostadinova, K. A.; Spears, R. J.; Chudasama, V. Tyrosine Bioconjugation – an Emergent Alternative. *Org. Biomol. Chem.* **2020**, 18, 9018.

(10) (a) Long, T.; Liu, L.; Tao, Y.; Zhang, W.; Quan, J.; Zheng, J.; Hegemann, J. D.; Uesugi, M.; Yao, W.; Tian, H.; Wang, H. Light-Controlled Tyrosine Nitration of Proteins. *Angew. Chem., Int. Ed.* **2021**, 60, 13414. (b) Wang, P.; Cheng, Y.; Wu, C.; Zhou, Y.; Cheng, Z.; Li, H.; Wang, R.; Su, W.; Fang, L. Tyrosine-Specific Modification via a Dearomatization-Rearomatization Strategy: Access to Azobenzene Functionalized Peptides. *Org. Lett.* **2021**, 23, 4137. (c) San Segundo, M.; Correa, A. Site-Selective Aqueous C-H Acylation of Tyrosine-Containing Oligopeptides with Aldehydes. *Chem. Sci.* **2020**, *11*, 11531. (d) Hu, Q.-L.; Hou, K.-Q.; Li, J.; Ge, Y.; Song, Z.-D.; Chan, A. S. C.; Xiong, X.-F. Silanol: A Bifunctional Group for Peptide Synthesis and Late-Stage Functionalization. *Chem. Sci.* **2020**, *11*, 6070.

(11) (a) Chudasama, V.; Maruani, A.; Caddick, S. Recent Advances in the Construction of Antibody–Drug Conjugates. *Nat. Chem.* **2016**, *8*, 114. (b) Bandyopadhyay, A.; McCarthy, K. A.; Kelly, M. A.; Gao, J. Targeting Bacteria via Iminoboronate Chemistry of Amine-Presenting Lipids. *Nat. Commun.* **2015**, *6*, 6561 and references cited therein. (12) Yang, Y.-Y.; Ascano, J. M.; Hang, H. C. Bioorthogonal Chemical Reporters for Monitoring Protein Acetylation. *J. Am. Chem. Soc.* **2010**, *132*, 3640.

(13) (a) Schneider, T.; Martin, J.; Durkin, P. M.; Kubyshkin, V.; Budisa, N. The Regioselective Synthesis of o-Nitrobenzyl DOPA Derivatives. *Synthesis* **2017**, *49*, 2691. (b) Chen; Zhu, Y.-F.; Wilcoxen, K. An Improved Synthesis of Selectively Protected L-Dopa Derivatives from L-Tyrosine. *J. Org. Chem.* **2000**, *65*, 2574.

(14) (a) Andrade-Sampedro, P.; Matxain, J. M.; Correa, A. Pd-Catalyzed $C(sp^2)$ -H Alkoxycarbonylation of Phenethyl- and Benzylamines with Chloroformates as CO Surrogates. *Chem. - Eur. J.* **2021**, 27, 5782. (b) Guerrero, I.; Correa, A. Cu-Catalyzed Site-Selective $C(sp^2)$ -H Radical Trifluoromethylation of Tryptophan-Containing Peptides. *Org. Lett.* **2020**, *22*, 1754. (c) San Segundo, M.; Correa, A. Pd-Catalyzed Site-Selective $C(sp^2)$ -H Radical Acylation of Phenylalanine Containing Peptides with Aldehydes. *Chem. Sci.* **2019**, *10*, 8872. (d) San Segundo, M.; Correa, A. Site-Selective Cu-Catalyzed Alkylation of α -Amino Acids and Peptides toward the Assembly of Quaternary Centers. *ChemSusChem* **2018**, *11*, 3893.

(15) (a) Enthaler, S.; Company, A. Palladium-Catalysed Hydroxylation and Alkoxylation. *Chem. Soc. Rev.* **2011**, *40*, 4912. (b) Shi, S.; Kuang, C. Palladium-Catalyzed Ortho-Alkoxylation of 2-Aryl-1,2,3-triazoles. *J. Org. Chem.* **2014**, *79*, 6105. (c) Zhang, C.; Sun, P. Palladium-Catalyzed Direct $C(sp^2)$ -H Alkoxylation of 2-Aryloxypyridines Using 2-Pyridyloxyl as the Directing Group. *J. Org. Chem.* **2014**, *79*, 8457. (d) Jiang, T.-S.; Wang, G.-W. Palladium-Catalyzed Ortho-Alkoxylation of Anilides via C-H Activation. *J. Org. Chem.* **2012**, *77*, 9504.

(16) (a) Dutta, U.; Maiti, S.; Bhattacharya, T.; Maiti, D. Arene diversification through distal $C(sp^2)$ -H functionalization. *Science* **2021**, *372*, eabd5992. (b) Rej, S.; Das, A.; Chatani, N. Strategic Evolution in Transition Metal-Catalyzed Directed C-H bond Activation and Future Directions. *Coord. Chem. Rev.* **2021**, *431*, 213683. (c) Sambiagio, C.; Schönbauer, D.; Blieck, R.; Dao-Huy, T.; Pototschnig, G.; Schaaf, P.; Wiesinger, T.; Zia, T. F.; Wencel-Delord, J.; Besset, T.; Maes, B. U. W.; Schnürch, M. A Comprehensive Overview of Directing Groups Applied in Metal-Catalysed C-H Functionalisation Chemistry. *Chem. Soc. Rev.* **2018**, *47*, 6603.

(17) (a) Lou, S.-J.; Chen, Q.; Wang, Y.-F.; Xu, D.-Q.; Du, X.-H.; He, J.-Q.; Mao, Y.-J.; Xu, Z.-Y. Selective C–H Bond Fluorination of Phenols with a Removable Directing Group: Late-Stage Fluorination of 2-Phenoxyl Nicotinate Derivatives. ACS Catal. **2015**, *5*, 2846. (b) Chu, J.-H.; Chen, S.-T.; Chiang, M.-F.; Wu, M.-J. Palladium-Catalyzed Direct Ortho Aroylation of 2-Phenoxypyridines with Aldehydes and Catalytic Mechanism Investigation. Organometallics **2015**, *34*, 953. (c) Liu, B.; Jiang, H.-Z.; Shi, B.-F. Palladium-Catalyzed Oxidative Olefination of Phenols Bearing Removable Directing Groups under Molecular Oxygen. J. Org. Chem. **2014**, *79*, 1521. (d) Yao, J.; Feng, R.; Wu, Z.; Liu, Z.; Zhang, Y. Palladium-Catalyzed Decarboxylative Coupling of α - Oxocarboxylic Acids with C(sp²)–H of 2-Aryloxypyridines. Adv. Synth. Catal. **2013**, 355, 1517.

(18) For more details, see the Supporting Information.

(19) (a) Guin, S.; Rout, S. K.; Banerjee, A.; Nandi, S.; Patel, B. K. Four Tandem C-H Activations: A Sequential C-C and C-O Bond Making via a Pd-Catalyzed Cross Dehydrogenative Coupling (CDC) Approach. Org. Lett. 2012, 14, 5294. (b) Yin, Z.; Sun, P. Palladium-Catalyzed Direct ortho-Acylation through an Oxidative Coupling of Acetanilides with Toluene Derivatives. J. Org. Chem. 2012, 77, 11339. (20) (a) Park, J.; Kim, A.; Sharma, S.; Kim, M.; Park, E.; Jeon, Y.; Lee, Y.; Kwak, J. H.; Jung, Y. H.; Kim, I. S. Direct Acylation of N-Benzyltriflamides from the Alcohol Oxidation Level via Palladium-Catalyzed C-H Bond Activation. Org. Biomol. Chem. 2013, 11, 2766. (b) Kim, M.; Sharma, S.; Park, J.; Kim, M.; Choi, Y.; Jeon, Y.; Kwak, J. H.; Kim, I. S. Pd-Catalyzed Oxidative Acylation of 2-Phenoxypyridines with Alcohols via C-H Bond Activation. Tetrahedron 2013, 69, 6552. (c) Xiao, F.; Shuai, Q.; Zhao, F.; Baslé, O.; Deng, G.; Li, C.-J. Palladium-Catalyzed Oxidative sp² C-H Bond Acylation with Alcohols. Org. Lett. 2011, 13, 1614.

(21) (a) Kumar, P.; Dutta, S.; Kumar, S.; Bahadur, V.; Van der Eycken, E. V.; Vimaleswaran, K. S.; Parmar, V. S.; Singh, B. K. Aldehydes: Magnificent Acyl Equivalents for Direct Acylation. *Org. Biomol. Chem.* **2020**, *18*, 7987. (b) Santiago, C.; Sotomayor, N.; Lete, E. Pd(II)-Catalyzed C-H Acylation of (Hetero)arenes-Recent Advances. *Molecules* **2020**, *25*, 3247.

(22) (a) Dobereiner, G. E.; Crabtree, R. H. Dehydrogenation as a Substrate-Activating Strategy in Homogeneous Transition-Metal Catalysis. *Chem. Rev.* 2010, 110, 681. (b) Guillena, G.; Ramón, D. J.; Yus, M. Alcohols as Electrophiles in C-C Bond-Forming Reactions: The Hydrogen Autotransfer Process. *Angew. Chem., Int. Ed.* 2007, 46, 2358.