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REVIEW

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Hexafluoroisopropanol: the magical solvent for Pdcatalyzed C-H activation

Among numerous solvents available for chemical transformations, 1,1,1,3,3,3-hexafluoro-2-propanol (popularly known as HFIP) has attracted enough attention of the scientific community in recent years. Several unique features of HFIP compared to its non-fluoro analogue isopropanol have helped this solvent to make a difference in various subdomains of organic chemistry. One such area is transition metal-catalyzed C-H bond functionalization reactions. While, on one side, HFIP is emerging as a green and sustainable deep eutectic solvent (DES), on the other side, a major proportion of Pd-catalyzed C-H functionalization is heavily relying on this solvent. In particular, for distal aromatic C-H functionalizations, the exceptional impact of HFIP to elevate the yield and selectivity has made this solvent irreplaceable. Recent research studies have also highlighted the Hbond-donating ability of HFIP to enhance the chiral induction in Pd-catalyzed atroposelective C-H activation. This perspective aims to portray different shades of HFIP as a magical solvent in Pdcatalyzed C-H functionalization reactions.

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Introduction

The crucial role of a solvent in governing the pathway of a chemical reaction was realized long ago when the great Greek philosopher Aristotle stated that "...compounds do not react unless fluid or if dissolved".1 Although solid-state reactions have been well established in recent times, they are bound to certain terms and conditions. A solvent can directly influence stereochemistry of the intermediate, and hence the product's (recalling the role of solvents in nucleophilic substitution reactions). All of these phenomena are controlled by a few major parameters, which include acidity, ionic strength, boiling point, H-bonding, or other weak interactions, along with supplementary physical properties. While there are plenty of solvents known for their extraordinary assistance in various synthetic transformations, there are a few hidden gems often tagged as "unconventional solvents".2 1,1,1,3,3,3-Hexafluoro-2-propanol or HFIP is one of them. It could be the corrosive nature (highly burning sensation upon inhalation or skin contact) or inflammability of HFIP that mandates extra care while handling

the rate of a reaction, the formation of a particular product, the

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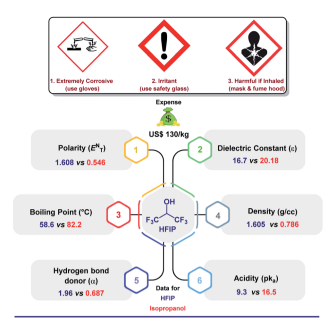


Fig. 1 A comparative overview of different physical characteristics, price, and safety details of HFIP and isopropanol.^{20,21}

and storage or being expensive (US\$130 per kg)3 that holds back researchers across the globe to use the solvent immensely. Despite this quandary, HFIP has earned its eminence in various research domains due to its spectacular physical and chemical properties. Fig. 1 features a comparative statistics of the solvent with isopropanol comprising minute to drastic differences in physical properties, which eventually dictates the mode of a chemical reaction. Unlike other analogues, the presence of two tri-fluoroalkyl groups in HFIP is presumed to remarkably alter the course of a chemical reaction and leverages great opportunities in traditional organic synthesis, electrocatalysis, photocatalysis, biological studies, and even in environmental science. Before we dive into the main part, we take this prospect to describe glimpses of different underprivileged applications of HFIP concisely to highlight its widespread acceptability in various research fields.



Debabrata Maiti received his PhD from Johns Hopkins University (USA) in 2008 under the supervision of Prof. Kenneth D. Karlin. After postdoctoral studies at Massachusetts Institute of Technology (MIT) with Prof. Stephen L. Buchwald (2008–2010), he joined the Department of Chemistry at IIT Bombay in 2011, wherein he is currently an Associate Professor. His research interests focus on the develop-

ment of new and sustainable catalytic methods.

1.1 Traditional organic synthesis

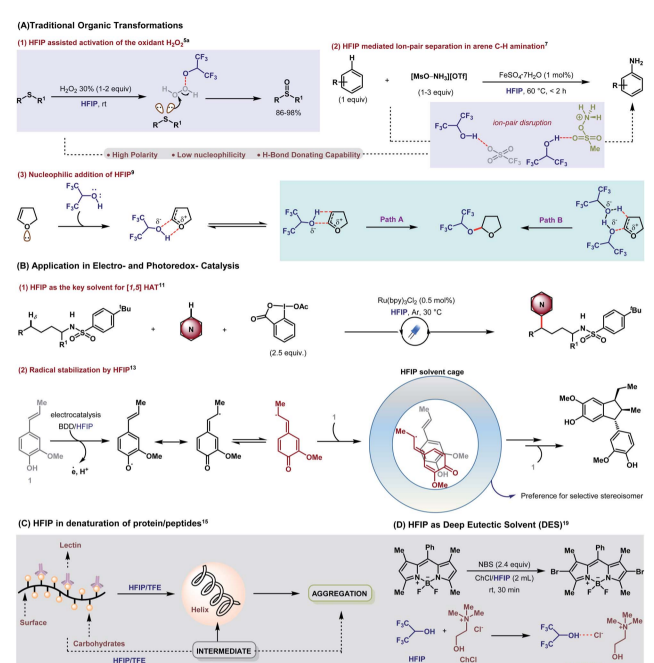
With the incorporation of fluorine, (i) HFIP becomes a befitting excellent H-bond donor and (ii) the polarity of the O-H bond is enhanced, which eventually increases the acidity. In a recent chemical record, Xiao and An have precisely displayed the impact of such substantial properties of HFIP in different genres of synthetic organic chemistry.4 By exploiting the H-bonding capability,5 the solvent can activate key oxidizing agents in extremely useful oxidation reactions (e.g., H2O2 in sulfide to sulfoxide,5a Scheme 1A, eqn (1), Baeyer-Villiger oxidation of ketone and epoxidation of olefins5b,c); in other instances such as in a faster Diels-Alder reaction by stimulating the dienophile, 6a enabling cyclopropane ring-opening reactions^{6b} and other useful organic transformations.6c In a recent report, Ritter group explains that Hbonding exerted by HFIP often leads to ion-pair separation which indeed promotes [MsO-NH3](OTf), the precursor for arene C-H amination (Scheme 1A, eqn (2)).7 Being non-nucleophilic, the solvent also promotes site-specific ring opening of 1-CF3substituted epoxy ethers.8 While synthetic chemists often cherish the non-nucleophilic nature of HFIP in different aspects, there are instances where HFIP takes part in nucleophilic addition. One such example is demonstrated by Bonnet-Delpon where a hexafluoropropyloxy acetal can be availed in synthetically useful yield in a HFIP-assisted ion-pair interaction (Scheme 1A, eqn (3)).9

1.2 En route to green chemistry: electro- and photoredox catalysis

The remarkable solvent effect of HFIP is not constrained only to the conventional synthetic platform, rather it is quite impressing in electrocatalysis as well as photocatalysis. Excellent redox stability in combination with a high dielectric constant (ε , 15.7) makes HFIP perfect for photoredox chemistry. The superpolar nature of the solvent [10² factor higher than trifluoroacetic acid (TFA) and by a factor of $\sim 10^8$ compared to acetonitrile (MeCN)] assists in stabilizing radical cations, which are a frequent intermediate in electro- and photoredox catalysis. 10 Probably, due to the same reason, the solvent is found to be way more efficient than any other polar solvents in a recent δ -C-H heteroarylation of protected aliphatic amines via [1,5]HAT (Scheme 1B, eqn (1)).11 Because of the high reactivity of the radical cations, the whole catalysis predominantly depends on the reaction media. It has been observed that non-nucleophilic HFIP can easily solvate the counter anions by H-bonding interactions, thus leaving the radical cations free.12 Recently, Boron Doped Diamond (BDD)-supported electrodes in HFIP medium have emerged as a proficient tool in electrocatalysis. In anodic oxidation of isoeugenol in a 1,1,1,3,3,3hexafluoro-2-propanol (HFIP)/(BDD) electrode system, HFIP forms a H-bonding network to surround the key radical intermediate. Inside the solvent cage, the intermediary radical species has a specific orientation to attain the highest stereoselectivity (Scheme 1B, eqn (2)).13

1.3 Biological experiments

Apart from typical synthetic chemical transformations, HFIP is very common in the hands of biologists. The effects of alcoholic



Scheme 1 Exposure of 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP) in different domains of experimental science.

solvents on proteins and peptides, to find out the cause of their stability inside the biological system, are a topic of great importance. Seminal studies indicate two major causes of alcohol-mediated protein denaturation (a) disruption of the natural state and (b) introduction of an α -helical motif. Polar and protic alcoholic solvents disrupt the distal hydrophobic interactions and elevate the local polar or hydrophilic interactions via weak interactions such as H-bonding. It is worth mentioning that HFIP with six fluorine atoms significantly induces α -helical conformation and imposes local polar interactions to a higher extent, which leads to protein denaturation.

Additionally, hexafluoroisopropanol indirectly influences the structural modification of protein and lipid layers of different biological membranes (Scheme 1C).¹⁵ Apart from peptide chemistry, HFIP is potential enough to produce lipid bilayer leakage and alter the biochemical properties of the lipid phase.¹⁶

1.4 Deep eutectic solvents

(150 mmol)

(100 mmol)

Traditional volatile organic compounds (VOCs) as solvents exert severe adverse impact on the environment such as easy accumulation in the atmosphere (due to low boiling point) causing

ChCI/HFIF

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respiratory trauma, flammability, high-grade toxicity, and most importantly, non-biodegradability.17 Water as a cheap and abundant solvent could be a probable alternative; however, lack of solubility of organic reagents does not meet the benchmark. In this prospect, deep eutectic solvents (DESs) have come up as an impending option to balance the nature and human innovations. A DES generally consists of a eutectic mixture of two or three species capable of self-aggregation where at least one of them is a hydrogen-bond donor (HBD) and the other is a hydrogen-bond acceptor (HBA). Although chlorine-based solvents (e.g., chloroform, tetrachloromethane, etc.) and even imidazole-based ionic liquids (ILs) are well recognized as leading components of DESs, being expensive and having noxious nature often raise severe threat to the environment.18 To bypass such hindrances, 1,1,1,3,3,3-hexafluoro-2-propanol as a HBD is evolving as a green and sustainable DES. HFIPbased aqueous biphasic solvents (ABSs) are now extensively used for the microextraction of components from various beverages, partitioning of dyes, as the green medium for diverse organic synthesis, recycling of food wastes, separation of natural products and so on.19

Over years, several reviews20 have come up to render the bigger picture of HFIP in many domains albeit focused reviews on the direct implication of HFIP in transition metal-catalyzed C-H bond functionalization are very scarce.²¹ It is worth specifying that the reliability of the solvent with palladium catalysts has taken the C-H bond transformations to a greater extent. The scope of this perspective is to highlight an irreplaceable plea for HFIP in Pd-catalyzed C-H activation. As our primary interest, we have emphasized on "how" HFIP is affecting the nature of a reaction. In particular, we describe the direct influence of a weak interaction force like H-bonding to stabilize the metallacycles and/or to interrelate with a suitable substrate to promote stereo- and regioselectivity. While the prime focus of this review is Pd-catalyzed C-H activation reactions, the intimacy of HFIP is not constrained only to Pd, rather its compatibility with other transition metals is also well inculcated in literature.22 For obtaining a better insight, we have subclassified the entire topic into four major parts commencing with proximal C-H activation.

2. Impact of HFIP in Pd-catalyzed proximal C-H functionalizations

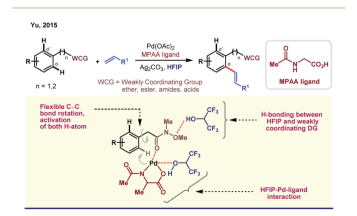
It has been a century since C–H bond functionalizations or precisely C–H bond activation has added a new dimension to contemporary organic synthesis. The amplitude of C–H bonds in natural products and pharmaceuticals has compelled the global synthetic chemists to strategize site-selective C–H functionalization as a transformative paradigm to access such valuable motifs. In this section on the perspective, we discuss the influence of HFIP on Pd-catalyzed regiospecific proximal C–H activation reactions (including ortho-C(sp²)–H and α/β -C(sp³)–H bonds). Evidently, transition metal-catalyzed ortho-C(sp²)–H functionalizations are well flourished because of the thermodynamically formed five- or six-membered

metallacycles. Since proximal C–H activation is nearly a favorable situation, it occasionally needs any further driving force. Hence, very few cases are known where HFIP is highly essential to attain proximal selectivity or excellent yield.

2.1 C(sp²)-H functionalizations

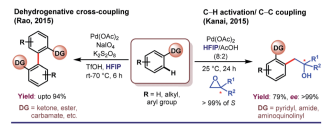
One of the central roles of HFIP in proximal aromatic C-H activation is stimulation of the weakly coordinating directing groups (WCGs) by H-bonding. Intrigued by this observation, Yu and coworkers demonstrated an ether-directed ortho-olefination of aromatic substituents in 2013.23a The strategy was extended thereafter to diolefination and acetoxylation by exploring other weakly coordinating groups such as Weinreb amides, ketones and esters (Scheme 2).23b Enantioselective iodination and acetoxylation of mandelic acid were also reported by the same group using HFIP as the principal solvent.24 Interestingly, HFIP was found to be the appropriate solvent when the reaction followed the Pd(II)/Pd(IV) pathway, whereas tert-amyl-OH appeared to be the best for the Pd(0)/Pd(II) cycle. However, readers should keep in mind that the catalytic cycle may vary from case to case, and HFIP is not always the preeminent choice of solvent in every Pd(II)/Pd(IV) cycle.25

HFIP also serves as the chief solvent in different oxidant-free ortho-functionalizations under mild reaction conditions. Kanai and co-workers reported C-H activation, followed by coupling of oxiranes with 2-phenyl pyridine derivatives to generate orthoalkylated compounds (Scheme 3).26 This protocol is exceptionally proficient at room temperature and acquires stereoretention with 99% ee in the product. The first report on weak coordination-promoted Pd-catalyzed dehydrogenative crosscoupling reaction of ketones, esters, and carbamates was made by Rao group in HFIP medium (Scheme 3).27 Later on, Su and co-workers reported the arylation of electron-deficient benzoic acid,28a and recently, Li group has also reported a ligand-controlled ortho-arylation of 2-phenylethylamines in HFIP.28b A recent work by Li and Wang shows the remarkable solvent effect by HFIP on a Pd(II)-catalyzed ortho-olefination of arenes directed by a polyfluoroalkylsulfinyl auxiliary.28c In this context, it is worth mentioning that several seminal reports on



Scheme 2 HFIP favoring proximal *ortho-*C(sp²)–H olefination of weakly coordinating directing groups.

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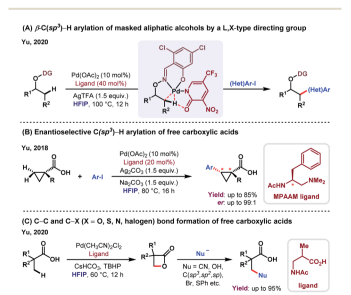
Scheme 3 Pd-catalyzed C(sp²)-H bond functionalization reactions.

 $Pd(\pi)$ -catalyzed *ortho*-C-H functionalizations in recent times have actually evolved employing HFIP as the supreme solvent.²⁹

2.2 C(sp³)-H functionalizations

The unique activity of the weakly tethering groups in HFIP can be witnessed in the case of aliphatic $\beta\text{-C}(sp^3)\text{-H}$ arylation of $\alpha\text{-amino}$ acids, 30 di-peptides 31 (both N-terminus and C-terminus) and Weinreb amides. 32 Excluding arylation, other functional transformations like olefination 33 and acetoxylation 34 of free carboxylic acids are quite sensitive toward HFIP. Other than acid derivatives, HFIP has turned out to be suitable for alcohols also. A recent report on methylene $\beta\text{-arylation}$ of masked aliphatic alcohol by the L, X-type auxiliary has again enlightened HFIP as the optimum solvent (Scheme 4A). 35 In 2018, MPAAM (mono-protected aminoethyl amine) ligand-assisted stereoselective $\beta\text{-C}(sp^3)\text{-H}$ arylation of aliphatic acid was also achieved by the same group with high enantioselectivity (up to 98% ee) in HFIP (Scheme 4B). 36

The extraordinary compatibility of HFIP with aliphatic acids gives leverage to chemists to unfold the other novel possibilities thereafter. For instance, β -lactonization of free acids, followed by ring-opening C-C and C-X (X = O, S, N, halogen, *etc.*) bond formation (Scheme 4C).³⁷ In both cases, HFIP was found to be



Scheme 4 Pd-Catalyzed β -C(sp 3)-H functionalization using HFIP as the key solvent.

the optimum solvent in $Pd(\pi)/Pd(\pi)$ catalysis. While the aforementioned reports justify like dissolves like theory for HFIP, seminal reports suggest the role of hexafluoro propanol in modulating the inert C-H bonds toward oxygenation.³⁸

On the other side, the influence of protic polar nature of HFIP was elegantly demonstrated by Larrosa group through a selective β-arylation method of thiophene and benzo[b]thiophene under mild reaction conditions.^{39a} The key intermediate of this Pd(0)/Pd(II) catalytic cycle was stabilized by H-bonding between two HFIP molecules. The simultaneous kinetic experiment and in silico studies revealed that the reaction follows Heck-type pathway (ΔG^{\ddagger} , 22.4 kcal mol⁻¹) instead of CMD (concerted metalation deprotonation) $(\Delta G^{\ddagger},$ and 24.7 kcal mol⁻¹, higher free energy barrier) as the latter process is thermodynamically formidable (Fig. 2). Even in an Agmediated α -arylation of benzo[b]thiophene by the same group, HFIP was used as the prime solvent.39h Along the lines, Xi,40a Li,40b Daugulis41 and Kuninobu42 have independently studied HFIP-promoted arylation of different aromatic and aliphatic systems.

3. HFIP as a key solvent for Pd-catalyzed distal C-H functionalizations

3.1 C(sp²)-H functionalizations

One fundamental challenge to execute site selectivity at the remote positions of arene is overcoming highly strained macrocyclic metallacycle formation. Although different strategies were employed to bypass this issue, most of them ended up with poor regioselectivity and limited substrate variety. HFIP, on the other hand, can stabilize the macrocyclic pre-transition state by its facile H-bond-donating capability. This extra solvent effect remarkably reduces the thermodynamic energy barrier for template-assisted distal C-H functionalizations. In this context, the first breakthrough was attained in 2012 by Yu group when they introduced a 'U-turned' weakly coordinating meta-selective nitrile template T1.43 The unique ability of HFIP to enhance the reaction outcome was observed for the metaolefination of hydrocinnamic acids (Scheme 5). Subsequently, the 'end-on' template was successfully employed for the metafunctionalizations of a series of different substrates like phenols, phenylacetic acids, phenyl propionic acids and Nheterocycles, where an excellent enhancement of the product yield, as well as meta-selectivity was observed in HFIP.44 In parallell, we have also introduced a novel and easy to install auxiliary T2 for meta-selective olefination of aryl acetic acids. However, trans-esterification of the substrate diminished the target product formation when trifluoroethanol was used as the solvent. Replacement of TFE by HFIP significantly suppressed the trans-esterification with improved yield and regioselectivity (Scheme 6).45 In other aspects, this simultaneous trans-esterification removed the template in situ without any further step.

On a different note, the catalytic amount of HFIP in DCE was enough to uplift the yield and selectivity for *meta*-olefination of arenes by a silyl-tethered nitrile motif by Tan and co-workers in 2013. ⁴⁶ Undoubtedly, apart from just being a solvent, HFIP was

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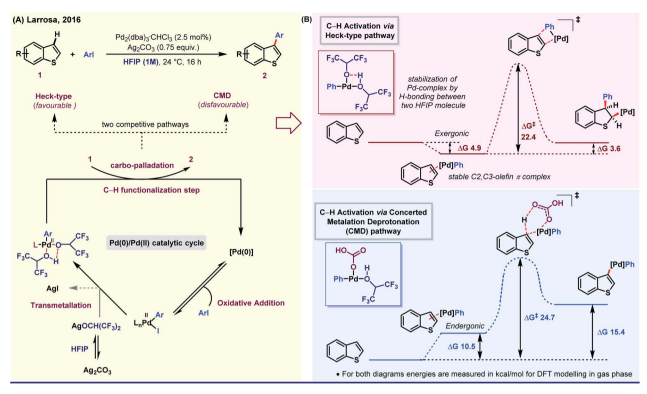
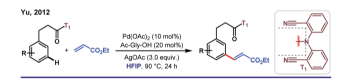


Fig. 2 (A) Pd-Catalyzed selective β -arylation of benzo[b]thiophene via a Heck-type pathway. (B) Energy profile diagram of two competitive pathways.



Scheme 5 First instance of Pd(\shortparallel)-catalyzed meta-C(sp 2)-H olefination of hydrocinnamic acids in HFIP medium.

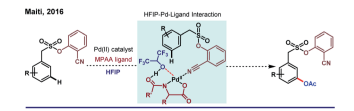


Scheme 6 Remarkable solvent effect by HFIP in Pd(II)-catalyzed meta- $C(sp^2)$ -H olefination of aryl acetic acids.

found to be extraordinary as an additive too. Stabilization of the electron-deficient metal center by HFIP is a well-known fact; however; experimental evidence of the fact for distal C-H functionalization is rare. Scheme 7 refers to HFIP and metal complex interaction, which is further confirmed by an NMR titration study during *meta*-acetoxylation of sulfonamides by our group.⁴⁷ It is worth mentioning that irrespective of modifications in coordination strength (from nitrile to pyrimidine),

HFIP has always played a remarkable supportive role to elevate the reaction outcome. Interestingly, the same legacy was followed when a carboxylate group was employed for the remote *meta*-selective olefination of hydrocinnamic amides by Li group in 2019. This work describes the predominance of κ^2 mode over κ^1 mode (which was more facile for proximal *ortho*-functionalization due to right conformational orientation) of the carboxyl group for any remote C–H functionalization for the very first time.

With ongoing surges in *meta*-C-H functionalizations, we thought to reach out to further distal C-H bonds. For the first time, the distal *para*-C-H bond of toluene was selectively functionalized by a silyl-tethered nitrile template of 'D' shape.⁵⁰ Building on seminal reports, we exploited the H-bond donor nature of HFIP to architect a better *para*-directing template. It was realized that a rotational restriction might orient the template toward the *para*-C-H bond completely to improve the *para*-selectivity (Fig. 3). A di-methoxy substitution was found



Scheme 7 HFIP-Pd-ligand interaction in Pd($_{\rm II}$)-catalyzed *meta-*C(sp²)-H acetoxylation of aryl sulfonamides.

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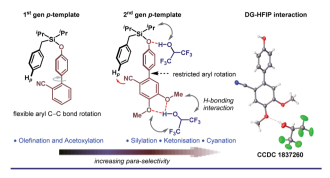


Fig. 3 HFIP-Substrate interaction in elevating para-selectivity.

optimal to induce such H-bonding with HFIP. This revised template (2nd generation template) indeed improved the *para*-selectivity drastically for other C–H to C–C/C–X functional transformations. Viability of the hypothesis was probed by NMR titration study, which reflected a significant substrate—HFIP interaction. However, it is not always the polarity or H-bond-donating ability, rather the exceedingly acidic character of HFIP also manipulates the mechanistic pathway of a reaction. For instance, when HFIP was substituted with its higher pK_a variants like d_2 -HFIP or isopropanol in the Pd-catalyzed p-ketonization reaction, no corresponding ketonized product was formed. This indeed proved the role of HFIP in protonating the vinyl ethers for generating a reactive intermediate for ketonization. These seminal insights made us even more

enthusiastic to explore other influences of HFIP in distal C-H activation. In our recent study, the crucial role of HFIP in Pdcatalyzed meta-allylation was monitored by NMR titration analysis.⁵² In contrast to other *meta*-selective templates, in T3, the pyrimidine moiety acts as a π -acceptor in addition to being a good σ-donor (Fig. 4A). Additionally, the weak H-bonding interaction between HFIP and pyrimidine 'N's as well as with 'O' in ether linker eventually makes the template statistically more approachable toward the meta-C-H bond. This study further ensures the stabilization of the key ion pairs by the polar protic nature of HFIP (Fig. 4B). A systematic NMR study clearly indicates a downfield shift of substrate protons, which further confirms the presence of an electronegative moiety, and thus, justifies the HFIP-substrate interaction and the role of the solvent (Fig. 4C). Very recently, our group has strategized to reach out the distal para-C-H bond of arene by Pd-norbornene (Pd/NBE) dual cooperative catalysis precluding the exact need of a para-selective template.53 With this, we report the first synthetic route to synthesize para-selective biaryls by C-H activation. Importantly, this strategy also involves HFIP as the sole solvent. The extraordinary effect of the solvent on the yield and regioselectivity is observed here as well. The compatibility of HFIP with Pd/NBE catalysis can also be seen in a recent work by Yu for meta-arylation of fluoroarenes.54

3.2 C(sp³)-H functionalizations

Although the role of HFIP in Pd(II)-catalyzed distal arene C-H functionalizations is acknowledged enough, it is a different

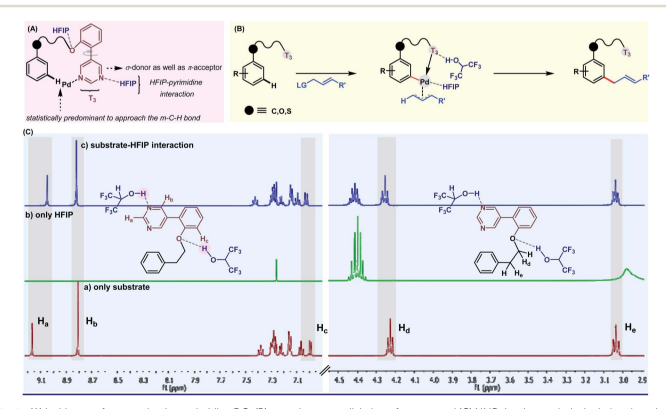


Fig. 4 (A) Inside out of *meta*-selective pyrimidine DG; (B) stepwise *meta*-allylation of arenes; and (C) NMR titration analysis depicting the role of HFIP in tuning substrate-DG orientation.

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Scheme 8 Distal γ -arylation of masked aliphatic alcohols.

story for aliphatic C-H activations where HFIP directly influences the reaction intermediates. One of the major problems associated with distal C-H functionalization is the involvement of biased metallation, which often limits its way to achieve homologous functionalizations.

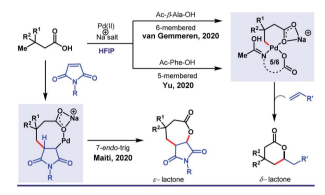
Distal γ-C-H functionalization of aliphatic alcohols demands the formation of a six-membered metallacycle over the thermodynamically accessible five-membered ring. In 2019, Yu and co-workers proposed a strategy for γ-arylation of aliphatic alcohols using 3-nitro-5-chloro-2-pyridone (L) as an external ligand via Pd(II)/Pd(IV) catalysis (Scheme 8).55 In this reaction, the alcohol was masked with an auxiliary T4 in such a way that the internal ring strain in the transition state allows [5,6]-fused bicyclic palladacycles over the [5,5]-bicylic one. The use of HFIP as the key solvent maintained the optimal solubility of the reagents and eventually stabilized the metallacyclic intermediate. In a subsequent work, the same group came up with γ-carbonylation of aliphatic alcohols in the form of a hemilabile benzyl ether.⁵⁶ While one end of the ether linker conceals the hydroxyl group of alcohol, the other end is linked with a bidentate ligand T5 which orients the palladium toward the target γ -C-H bond.

Upon migratory insertion of CO in int A, HFIP replaces the ligand from the metal coordination site. Eventually, the carbonylated product is obtained in the form of hexafluoroisopropanoate ester (Scheme 9). Nevertheless, chelationassisted distal C(sp3)-H functionalizations require preinstallation and post-synthetic removal of the directing template, which make the entire protocol sluggish and stepconsuming. Alternatively, directing the ability of native functional groups has remained restricted to the proximal site.

The reaction is well compatible with a wide range of aryl coupling partners and exceptionally sensitive toward the nature of the solvent. Apart from the high altitude of solubility in HFIP, the protic nature of the solvent plays a significant role in maintaining the optimal pH of the reaction medium.57

Scheme 9 HFIP stabilizing intermediate palladacycle for γ -carbonylation of aliphatic alcohols

Scheme 10 Pd(II)-catalyzed γ-arylation of free carboxylic acids in **HFIP**



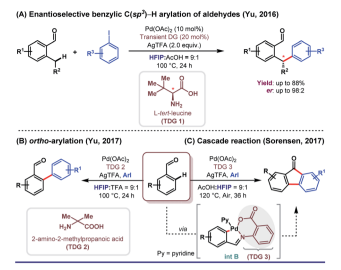
Scheme 11 HFIP as the sole solvent in Pd(\parallel)-catalyzed γ -olefination of free carboxylic acids to generate six-membered lactones.

Recently, van Gemmeren⁵⁸ and Yu⁵⁹ group independently extended the scope of the protocol. Unlike arylation, γ -olefination demands an external ligand, which is often a mono-Nprotected amino acid (Scheme 10). Indeed, in this case also, HFIP attributes for the highest experimental efficiency. While two different groups highlighted different aliphatic acids separately, our group simultaneously discovered an unusual seven-membered ε-lactone formation by utilizing maleimide as an unconventional source of olefin (Scheme 11).60

HFIP in Pd-catalyzed transient directing group-assisted C-H **functionalizations**

HFIP as a solvent and/or a co-solvent has played an immense role in different aspects of directing group (DG)-assisted C-H functionalization, as we have already mentioned earlier. However, transient directing group (TDG)-assisted C-H functionalization has emerged as a promising approach to circumvent the existing complications associated with the DG approach. TDGs have added a new dimension for synthetic organic chemists because of its several advantages. First, only a catalytic amount of use of TDGs; second, being economical and less tedious i.e., installation of TDGs, functionalization of C(sp²/sp³)-H bonds, and removal of TDGs, all were done in a single-pot reaction. 61,62 Like other sub-domains of C-H activation reactions, HFIP is equally effective as a solvent and sometimes as a co-solvent in a cocktail of solvents (a mixture of two or three solvents with different proportions) to execute TDG-assisted C-H activation. However, the actual role of HFIP in different Pd-catalyzed transient auxiliary mediated C-H activation has not been well discussed significantly.

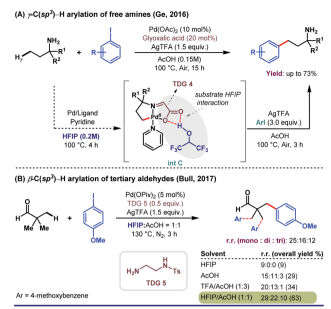
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Scheme 12 Different roles of HFIP in combination with Pd catalysis using transient auxiliaries (A) and (B) as the major solvent and (C) as additives in ortho-arylation of benzaldehydes.

In this context, the first example of using HFIP was put forth by Yu and co-workers in 2016 where they described enantioselective benzylic C(sp³)-H arylation of aldehydes enabled by a chiral TDG 1 L-tert-leucine in HFIP (Scheme 12A).62 A better productivity (up to 98:2 er) was obtained when a catalytic amount of AcOH was added in HFIP (HFIP: AcOH = 9:1, v/v). Recently, the same group has demonstrated an enantioselective β-C(sp³)-H arylation of cyclobutyl ketones utilizing a similar concept.63 Enhanced acidity was the foremost requirement to pursue the functionalization using different TDGs, thereby delivering the best possible result. Recently, Gou et al. depicted site-selective C(sp³)-H arylation of phenylacetaldehydes centered on a similar idea.⁶⁴ Later, in 2017, the same group reported ortho-C(sp²)-H arylation of substituted benzaldehydes, where HFIP was used as the key solvent (Scheme 12B).65 α-Substituted amino acids were found to be extremely proficient as a transient moderator (TDG 2) for such transformation. With the same perception, Sorensen and co-workers explored the synthesis of substituted fluorenones from benzaldehydes and aryl iodide through the Pd(II)-catalyzed C(sp²)-H functionalization reaction (Scheme 12C).66 However, here authors found 2amino benzoic acid (TDG 3) as the most promising transient auxiliary. Interestingly, HFIP played a prominent role in the isolation of [5,6]-fused palladacycle (int B) albeit being used as an additive (Scheme 12C).

Besides aromatic aldehydes, HFIP is also well attuned with aliphatic aldehydes, ketones, and primary amines. The polar protic nature of HFIP immensely manipulates the solubility of the substrates apart from maintaining the optimum pH of reaction media. In 2016, Ge and co-workers reported the β-C(sp³)-H arylation of aliphatic aldehyde enabled by 3-aminopropionic acid (TDG 4).67 The cocktail of HFIP and AcOH in a volumetric ratio of 5:1 showed a dramatic improvement in yield (from 29% to 75%) compared to that of only HFIP. Another interesting result of site-selective $C(sp^3)$ -H γ -arylation of



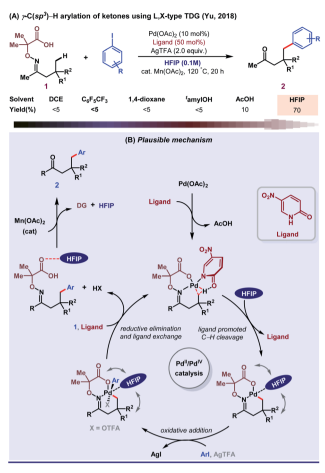
Scheme 13 Pd-catalyzed TDG-assisted C(sp³) – H arylation. (A) HFIP as an optimum solvent in the synthesis of intermediate; (B) solvent effect on the overall yield and selectivity.

primary aliphatic amines was published by the same research group employing glyoxylic acid as a TDG (Scheme 14A).68 They discovered AcOH as the optimal solvent for arylation with better yield and excellent site selectivity, yet HFIP played an unprecedented role in isolating the palladacycle as previously observed.

Although the authors have not provided detailed computational and mechanistic studies regarding the exact role of HFIP in the formation of intermediate (int C), it is very likely that HFIP induces an H-bonding interaction with the 'O' center of glyoxylic acid TDG, which further accelerates the reaction in the forward direction. Later, Bull and co-workers reported the β-C(sp³)-H arylation of tertiary aldehydes, facilitated by a transient imine directing group (Scheme 13B).69 Only HFIP was ineffective for this reaction, where a 1:1 mixture of HFIP and AcOH provided the best result. This solvent combination has increased not only the overall yield but also the amount of mono-arylated products compared to di- and tri-arylated products.

Until 2018, aliphatic TDG-assisted C(sp³)-H functionalizations were mostly restricted to the β-position of free amines, ketones, and aldehydes. HFIP was suitable for the γ -C(sp³)-H (hetero)arylation of ketones using 2,2-dimethyl aminooxyacetic acid auxiliary enabled by 2-pyridone ligand (Scheme 14A).70 The role of HFIP is attributed to its H-bonding interaction with the substrate for the formation of a six-membered palladacycle intermediate and in the successful removal of DG. Mechanistic studies revealed how ligand exchanges its position with HFIP, followed by an oxidative addition to further accelerate the Pd(II)/ Pd(IV) catalytic cycle and stabilize the Pd(IV) intermediate (Scheme 14B). Second, they described a method of ligand directed selective γ- and δ-C(sp³)-H arylation of free amines by a bidentate imine and a carboxylate L, X-type directing group.71

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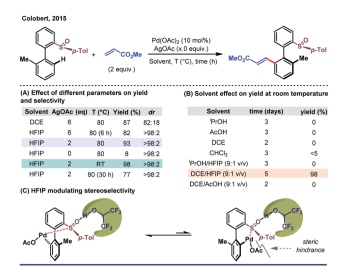
Scheme 14 (A) γ -C(sp³)-H arylation of ketones; (B) mechanistic blueprint displaying the detailed role of HFIP.

This result was the outcome of the cooperative effect between the transient directing group and external pyridone ligands in HFIP solvent medium. In addition, other research groups like Jin,⁷² Ge and Li,⁷³ and Chen in collaboration with He,⁷⁴ Wei,⁷⁵ and Zhang⁷⁶ also contributed with their valuable results to enrich this area. However, the exact role of HFIP in the aforementioned reactions is still unknown, presumably due to the lack of computational and mechanistic studies.

HFIP: boosting stereoselectivity in Pd-catalyzed asymmetric C-H activation

Irrespective of different modes of action (as a solvent or an additive or a chiral auxiliary), fluorinated alcohols (TFE and HFIP) have gained enough acceptance for their remarkable impact on elevating stereoselectivity. Before its direct influence in asymmetric C-H activation, the unprecedented impact of HFIP as a solvent in a Pd-catalyzed asymmetric hydrogenation reaction was realized by Uneyama's group long back.77

Despite its uninterrupted application in other areas of homogeneous catalysis,78 HFIP has just started its journey as



Scheme 15 Pd(II)-catalyzed atroposelective olefination of biaryls. (A) Overall optimization of the reaction; (B) effects of different solvents at room temperature on yield; and (C) steric induction by HFIP favoring one diastereoisomer

a game changer in stereoselective C-H functionalization.214 It is hypothesized that the weak coordination of fluorinated alcohols stabilizes the active catalyst. Moreover, being a strong H-bond donor, HFIP can easily manipulate its adjacent C-H bond or other H-bond acceptors to stabilize or destabilize a particular intermediate or substrate. Realizing the crucial role of HFIP in C-H activation, Colobert and Wencel-Delord re-discovered a Pdcatalyzed atroposelective olefination of a biaryl sulfoxide motif in HFIP solvent. The reaction previously performed in DCE led to the desired olefinated product with a compromised yield and enantioselectivity at the cost of superstochiometric use of AgOAc (6 equiv.) and elevated reaction temperature (Scheme 15A).⁷⁹ A dramatic improvement in yield and enantioselectivity was observed when HFIP was employed, whereas other polar and/or acidic solvents failed or took a longer time to deliver the desired product at room temperature (Scheme 15B). Based on several seminal reports, the authors suggested that the sulfoxide moiety present in the substrate is highly prone to Hbonding with HFIP which lengthens or alternatively weakens the S=O bond which alters the electronic environment of the sulfoxide group and consequently affects the rate of C-H activation step. Moreover, such H-bonding also adds an additional steric factor in the palladacycle, preferably allowing a less hindered atropoisomeric intermediate to react further (Scheme 15C). Another significant benefit of using HFIP as a solvent is reduction in excess use of AgOAc (6 to 2 equiv.) for the reaction, which could be possible by a HFIP facilitated re-oxidation of Pd(0) along with the Ag^I oxidant.

Subsequently, Shi and co-workers elegantly described the possibility of utilizing HFIP as a solvent in another Pd-catalyzed atroposelective olefination of bi-aryl aldehydes.80 In this work, a stark improvement in yield and enantioselectivity was observed when a 4:1 (v/v) combination of HFIP and acetic acid was employed (Scheme 16).

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vield (%)

Solvent effect on yield and selectivity

Solvent TFE HFIP MeOH AcOH HFIP/AcOH (4/1 v/v)

Scheme 16 Effect of HFIP in enhancing the reaction yield.

Although this observation indicates the necessity of a low pH reaction media to improve the reaction outcome, the authors did not specify the actual role of HFIP in this reaction. Based on a similar concept, the authors came up with several new atroposelective functionalization reactions with HFIP being the 'best suitable solvent'.* In addition to the aforementioned works, several asymmetric C–H activation reactions regardless of substrate diversity have been performed using HFIP as the solvent.* Thorough optimization led the authors to choose HFIP over other solvents for better sustainability. Nevertheless, the actual role of HFIP has remained uninvestigated in a majority of cases.

Conclusions and outlook

Over the centuries, solvents have manipulated the direction of different chemical conversions by their steric and electronic features. For any catalytic reaction, it is not the catalyst that always dictates the competence of the protocol; rather it also depends on the reaction media. While we discuss the solvent effect in different aspects of organic synthesis, we cannot overlook their influence on biological systems and environment. In that perspective, 1,1,1,3,3,3-hexafluoro-2-propanol or HFIP is emerging as a solvent of choice. Especially, due to its multidirectional application in transition metal-catalyzed C-H functionalizations, the urge for HFIP is almost inimitable now. Although HFIP is a renowned solvent, this perspective portrays other deprived roles of HFIP (e.g. as additive, catalyst, activator, etc.) synchronously. As of our primary focus, we have precisely discussed three foremost impacts of HFIP in Pd-catalyzed C-H activation reactions. First, as we move from proximal (ortho-) to distal (meta- and para-) positions of an arene, HFIP helps in harnessing the transition metal catalyst with the target C-H bond through H-bonding interactions. Moreover, being a strong H-bond donor, HFIP is able to introduce steric crowding in the transitional steps and uplift stereoselectivity in many asymmetric C-H activations. Second, the polar protic environs of HFIP stabilizes macrocyclic cyclophanes like metallacycles (especially when metals are in a high oxidation state), ion pairs and radical cations involved in proximal and distal C-H activation. Third, as an additive, HFIP lessens the pH of the media, which eventually helps in solubilizing acidic and/or polar

substrates (e.g., acids, free amines and alcohols). In addition, this review embraces every other subtle aspects of HFIP in Pd-catalyzed C-H activation reactions in detail along with their mechanistical intricacies.

Despite its excellence, a pervasive limitation of using HFIP as a sustainable reagent could be its low cost efficiency and corrosive nature. While an extra layer of cautiousness can diminish the second concern, the other issue is also addressable by recycling the solvent through distillation (low bp of HFIP, 58.6 °C). Nevertheless, very few works have highlighted a detailed study to investigate the actual role of HFIP (whether solvent, co-solvent and/or additives) in Pd-catalyzed C-H activation reactions. Hence, more experimental and computational analyses will further amplify the other unaddressed aspects of HFIP in palladium catalysis in near future.

Conflicts of interest

There are no conflicts to declare.

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