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Introduction

Aryl C-glycosides are important compounds encompassing structurally diverse natural products and drugs.¹ The distinctive feature of these molecules – *i.e.* the presence of a stable C–C bond connecting a saccharide unit to an aryl moiety – confers resistance towards hydrolytic enzymes, which makes them privileged glycomimetic candidates in drug development programs (Scheme 1a).² Given their relevance, numerous synthetic methods for the efficient preparation of aryl C-glycosides have been developed in recent years, most of them achieving the formation of the aryl C-glycoside linkage through cationic, anionic, and radical glycosyl intermediates, or with the use of transition metal catalysts.^{1b,3}

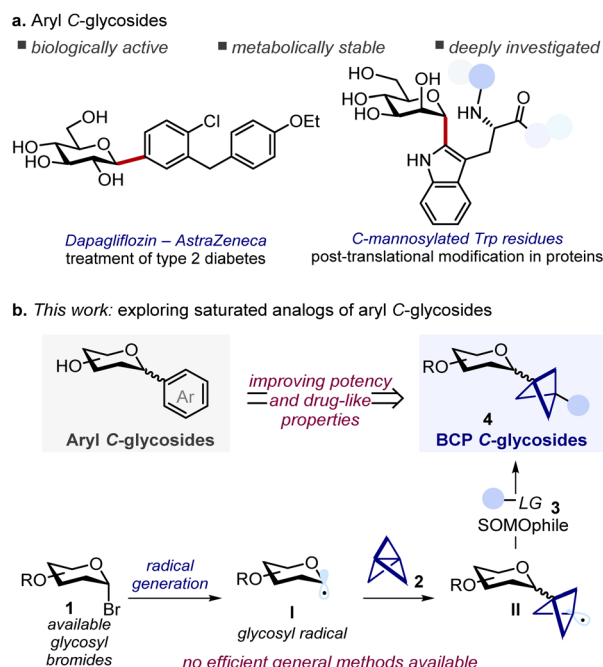
On the other hand, the replacement of planar conjugated moieties with three-dimensional fragments has been established as a potent strategy in medicinal chemistry to improve the physicochemical properties of lead compounds and increase their success rate in clinical trials.⁴ Driven by this principle, in the past few years synthetic chemists have focused on the development of C(sp³) rich bioisosteres able to replace planar aromatic cores, which are highly represented motifs in

A radical strategy to the synthesis of bicyclo[1.1.1]pentyl C-glycosides

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Aryl C-glycosides, in which carbohydrates are directly linked to aryl fragments through a stable C–C bond, are an important class of biologically active molecules widely found in nature. These compounds exhibit resistance to (enzymatic) hydrolysis, a property that has been successfully leveraged in the development of metabolically stable drugs. On the other hand, despite their potential, three-dimensional analogues of aryl C-glycosides remain largely overlooked. Here, we present a three-component radical strategy that grants access to this underexplored chemical space. Specifically, glycosyl bromides serve as a source of glycosyl radicals, which can react with [1.1.1]propellane and a suitable SOMOphile to afford bicyclopentyl C-glycosides. These C(sp³)-rich analogues replace a planar aryl ring with a three-dimensional bicyclopentyl moiety, which is expected to enhance physicochemical properties. The protocol is practical, mild, and amenable to scalable synthesis in continuous flow. Experimental and computational studies support a radical chain mechanism under kinetic control.

drugs.⁵ Among these, the saturated bicyclo[1.1.1]pentane (BCP) is arguably the most studied bioisostere of phenyl rings.⁶ When 1,3-disubstituted at the bridgehead positions, this core acts as



Scheme 1 (a) Representative examples of aryl C-glycosides as natural products and drugs. (b) Three-component radical approach towards bicyclo[1.1.1]pentyl (BCP) C-glycosides as saturated analogs. Trp: tryptophan; LG: leaving group; SOMO: singly occupied molecular orbital; BCP: bicyclo[1.1.1]pentyl.

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a rigid spacer projecting the substituents with exit vectors at approximately 180° that well mimic the spatial arrangement of *para*-disubstituted arenes. As such, bioisosteric replacement strategies with BCPs have been successfully applied in drug discovery programs, allowing the preparation of saturated analogs with unaltered targeting ability that show improved potency or enhanced physicochemical properties (*e.g.* solubility, and metabolic stability), and that can enable patent circumvention.⁷

Despite their potential, the development of saturated analogues of aryl *C*-glycosides has remained largely underdeveloped, which precludes the study of their properties and implementation in drug development campaigns. A possible explanation for this synthetic gap is the paucity of methods for the introduction of highly three-dimensional fragments at the anomeric position of saccharides. Indeed, while *C*-glycosylation strategies for the functionalization with alkyl, alkenyl, and alkynyl groups are well developed (either *via* polar or radical chemistry),^{3b,d,8} these approaches are mainly limited to the instalment of linear flexible carbon chains that do not have the requisites to properly mimicking the defined exit vectors provided by rigid aromatic scaffolds.

Restricting our focus to saturated *C*-glycosides featuring a BCP unit directly linked to the anomeric position, we envisioned that this moiety could be conveniently introduced by functionalization of [1.1.1]propellane (Scheme 1b).⁹ This strained precursor exhibits omniphilic reactivity, being able to react with electrophiles, strong nucleophiles, and radicals. Notably, radical addition to [1.1.1]propellane proceeds under remarkably mild conditions,¹⁰ making it particularly suited for functionalizing complex substrates such as glycosides.^{3d,11} In this regard, rare preparations of BCP *C*-glycosides have been independently reported by the Anderson and Molander groups by ring opening of [1.1.1]propellane with glycosyl radicals,^{10n,o} although both protocols involve the use of either unstable radical precursors (*e.g.* glycosyl iodides), transition metal-based photocatalysts, or expensive SOMOphiles (*e.g.* Sugimoto reagent). Notably, although these methods targeted a broad range of alkyl iodide and bromide substrates, their application to *C*-glycosyl derivatives was limited. Both reported just a single glucoside derivative and, in one case, with an unusual β -stereoselectivity that we believe misassigned. Furthermore, the scalability and further derivatization of the *C*-glycoside products were not investigated. With this work we aim to address these limitations providing a general versatile strategy to the efficient synthesis of BCP *C*-glycosides to enable their study and characterization.¹²

Taking inspiration from the work of the Molander group, we recognized that readily available glucosyl bromide **1** could serve as a radical precursor to glycosyl radical **I**. This reactive intermediate would then engage in a radical addition to [1.1.1]propellane **2** giving the BCP radical **II**.^{10n,o} Finally, reaction with a suitable SOMOphile **3** would deliver the desired BCP *C*-glycoside **4** (Scheme 1b). We reasoned that careful selection of the SOMOphile **3** would be key to the success of this three-component radical approach. Here, two key challenges must be considered: (i) glycosyl radical **I** has to discriminate between propellane **2** and SOMOphile **3**, leading to the selective

formation of the key intermediate **II**; (ii) **II** must then react selectively with SOMOphile **3** over propellane **2**, thus preventing staffane side-products. Herein we detail our efforts in implementing this strategy to provide general and efficient access to saturated aryl *C*-glycoside analogs. Two distinct protocols for the hydroglycosylation and carboboration of [1.1.1]propellane were developed, providing synthetically useful sugar-based building blocks that enabled smooth diversification into a small library of BCP *C*-glycoside derivatives. A thorough mechanistic investigation is also presented.

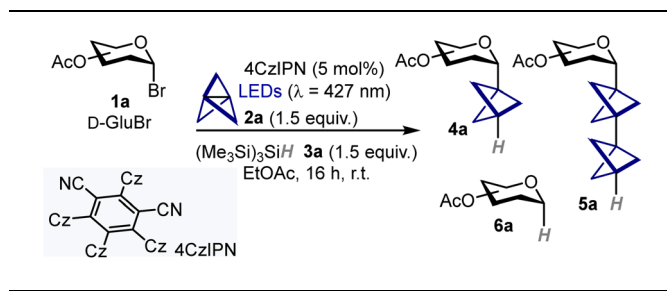
Results and discussion

Strategy validation – hydroglycosylation of propellanes

To test the feasibility of our approach, we began investigating the model reaction between acetobromo- α -D-glucose **1a** (0.1 mmol), [1.1.1]propellane **2a**, and tris(trimethylsilyl)silane **3a** (Table 1).¹³ After optimization (for details, see SI, Section C.1) we found that using a slight excess of both propellane **2a** and silane **3a** (1.5 equiv.), along with 4CzIPN as photocatalyst (PC)¹⁴ under blue LED ($\lambda_{\text{max}} = 427$ nm) irradiation provided BCP *C*-glucoside **4a** in good yield and stereoselectivity (71% yield; $\alpha:\beta$, 7 : 1) as judged by ¹H NMR analysis of the crude mixture (Table 1, entry 1), while minimizing the formation of staffane **5a** and 1-deoxyglucoside **6a** (9% and 20% yield, respectively). Notably, we found that the use of trialkylamines as hydrogen atom donors suppressed the formation of the reduced side-product **6a**, although product **4a** was obtained in diminished yield and with substantial formation of staffane **5a** (Table 1, entries 2,3). Since this latter side-product is particularly difficult to purify by flash column chromatography from the desired *C*-glycoside **4a**, we selected silane **3a** as the best hydrogen atom transfer (HAT) donor species. Stoichiometry was also found to be a key parameter, as increasing the excess of either propellane **2a** and silane **3a** led to decreased performance (Table 1, entries 4–6). Pleasingly, only a slight decrease in yield was observed when performing the reaction on a 0.2 mmol scale (63% yield, Table 1, entry 7). Finally, control experiments revealed that the reaction can also proceed in the absence of the PC and light, albeit with lower efficiency and increased side-products formation (Table 1, entries 8–10). These results indicate that a radical chain reaction can be thermally initiated, although the process is relatively inefficient and the use of 4CzIPN as photoinitiator is beneficial.

With the best conditions in hand, we sought to evaluate the applicability and generality of this three-component hydroglycosylation protocol (Table 2). First, we demonstrated that the protocol is scalable to a 2 mmol reaction under batch conditions, giving **4a** in 42% yield after isolation (336 mg). Then, we varied the protecting groups of D-glucosyl bromide showing that benzoyl and pivaloyl esters are also well tolerated (**4b–c**). To our delight, the reaction was amenable to the functionalization of several hexoses, pentoses, and deoxysugars in their pyranosidic form, enabling the preparation of BCP *C*-glycosides of D-galactose, D-mannose, D-lyxose, L-rhamnose, and L-fucose in moderate to good yields and excellent a stereoselectivity (**4d–h**). A fluorinated D-galactose derivative was found to be a competent



Table 1 Optimization of the reaction conditions^a

Entry	Deviation from standard conditions	4a ^b (%)	5a ^b (%)	6a ^b (%)
1	—	71	9	20
2	iPr ₂ EtN instead of (Me ₃ Si) ₃ SiH	33	28	—
3	Et ₃ N instead of (Me ₃ Si) ₃ SiH	37	17	—
4	3 equiv. of 2a	48	24	23
5	3 equiv. of 3a	53	11	36
6	3 equiv. of 2a and 3a	59	11	28
7	0.2 mmol	63	7	30
8	No 4CzIPN	44	8	45
9	No light	34	13	48
10	No PC, no light, 40 °C	40	18	21

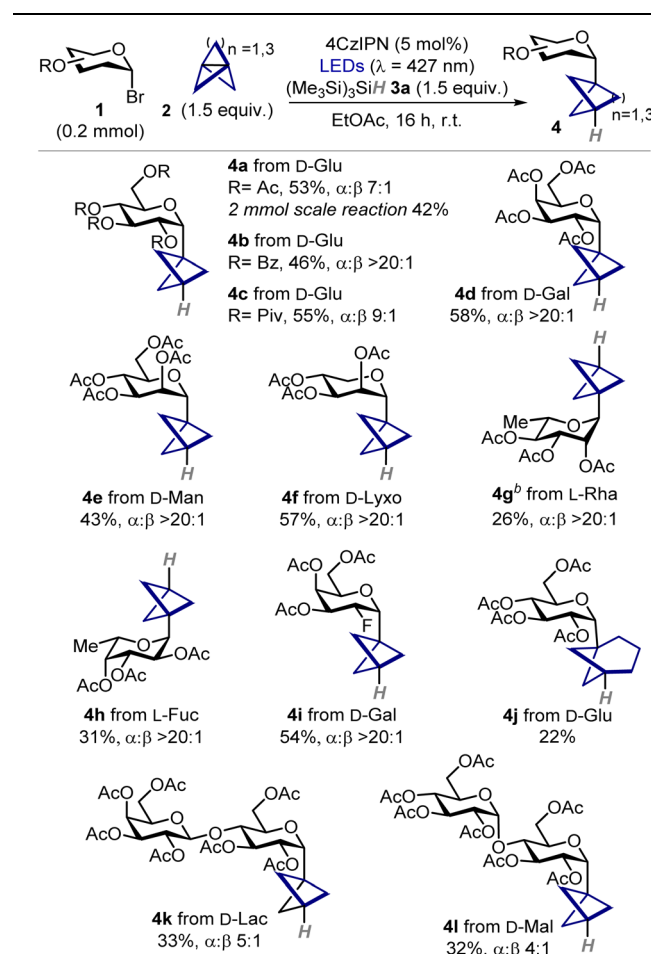
^a Reactions were performed using 0.1 mmol of 1a, [1.1.1]propellane 2a as solution in Et₂O, and 0.4 mL of EtOAc. ^b Determined by ¹H NMR analysis of the crude mixture using trichloroethylene as the internal standard. 4CzIPN: 1,2,3,5-tetrakis(carbazol-9-yl)-4,6-dicyanobenzene; Cz: carbazoyl; LED: light-emitting diode; r.t.: room temperature.

substrate (4i), showcasing the suitability of the method for the preparation of innovative chemical probes.¹⁵ When [3.1.1]propellane 2b was used we successfully obtained C-glycoside 4j whose bicycloheptyl moiety mimics a *meta*-disubstituted arene, albeit the reaction occurred in somewhat decreased yield.¹⁶ Finally, disaccharides bearing either a 1,4- α and 1,4- β glycosidic linkage gave the corresponding products 4k,l in synthetically useful yields. Remarkably, all products were generally obtained as the α anomers with very high levels of stereoselectivity. In all cases, BCP C-glycosides 4 were formed as the major products, along with traces of staffanes 5 and minor formation of 1-deoxyglycosides 6, as judged by ¹H NMR analysis of crude reaction mixtures.

Carboboration of [1.1.1]propellane towards BCP C-glycoside boronates

Next, we wondered whether this radical strategy could be expanded to other classes of SOMOphiles to access BCP C-glycosides of increased synthetic value. To our delight, after intense screening and optimization of key parameters we identified diboron(4) reagents as suitable reaction partners (for details see SI, Section C.2).^{10,17} Specifically, we found that glycosyl bromides 1 react in presence of an excess of [1.1.1]propellane 2a and bis(1,3-propanediolate)diboron (B₂pro₂) 3b to give BCP C-glycoside boronates 7 with good yield and good to excellent stereoselectivity (Table 3). Notably, this reaction occurs under thermal conditions – *i.e.*, in the absence of light or PC – with perfect selectivity for boronate 7. Since isolation by flash chromatography proved unsuccessful due to degradation

on silica gel, the Bpro moiety was instrumental, as it could be smoothly converted into the corresponding MIDA ester 8, enabling purification and full characterization.¹⁸ This protocol could be applied to both peracetylated and perbenzoylated glucosides (8a,b) as well as a variety of hexo- and pentopyranoses (8c–f) (Table 3). While these boronates were generally obtained with good to excellent stereoselectivity for the α anomer, a lower α : β ratio of 2:1 was observed for D-xylose derivative 8e, likely due to its higher conformational freedom.¹⁹ Deoxy sugars from the L-series also reacted well (8g,h). This enabled us to grow crystals suitable for X-ray analysis of the L-fucose derivative 8h, confirming the α configuration of the anomeric center and determining key structural parameters within the BCP moiety (see SI, Section E). Finally, we were able to successfully apply the protocol to a fluoro D-galacto derivative (8i) and to D-maltose and D-lactose disaccharides (8j,k).

Table 2 Light-driven hydroglycosylation of propellanes^a

^a Reactions performed on a 0.2 mmol scale using 4CzIPN (5 mol%), propellane 2 as solution in Et₂O (1.5 equiv.), and (Me₃Si)₃SiH 3a (1.5 equiv.). Yields refer to isolated products 4. α : β ratios were determined by ¹H NMR analysis of the crude mixture using trichloroethylene as the internal standard. ^b Reaction performed at 0.3 mmol scale. 4CzIPN: 1,2,3,5-tetrakis(carbazol-9-yl)-4,6-dicyanobenzene; LED: light-emitting diode; r.t.: room temperature.



Continuous flow scale-up and derivatization studies

To explore the synthetic potential of BCP *C*-glycosyl boronates, we scaled up the synthesis of glucoside **8a** and took advantage of the boronate group as a handle for further diversification.²⁰ Initial attempts to increase the reaction scale under batch conditions resulted in poor conversions due to slower kinetics. However, adapting the process to continuous flow using a homemade reactor enabled us to perform the reaction on a 2.0 mmol scale without loss of efficiency (see SI, Section F.2).²¹ Notably, the use of $B_2\text{pro}_2$ was crucial, as it ensured complete solubility of the reaction mixture, thereby facilitating a smooth transition to flow conditions. Finally, transesterification with MIDA afforded glucosyl boronate **8a** in 54% overall yield (Scheme 2).

While treating crude boronate **7** with oxone smoothly provides access to tertiary alcohol **9** (45% yield over two steps), MIDA ester **8a** could be efficiently converted into the corresponding tetrafluoroborate salt **10** (93% yield) (Scheme 2). Pleasingly, this key radical source participated in a Cu-catalyzed metallaphotoredox Chan-Lam reaction with 6-chloroindole, giving the corresponding C–N cross-coupled product **11**.¹⁰ Alternatively, **10** could also be employed in a Giese-type addition to dimethyl fumarate (**12**), and to methyl vinyl ketone when using a Cu(i) salt as a co-catalyst (**13**).²² Finally, a Ni-catalyzed metallaphotoredox protocol enabled the forging of a $\text{C}(\text{sp}^3)$ – $\text{C}(\text{sp}^2)$ bond, affording the cross-coupled product **14**.^{17d}

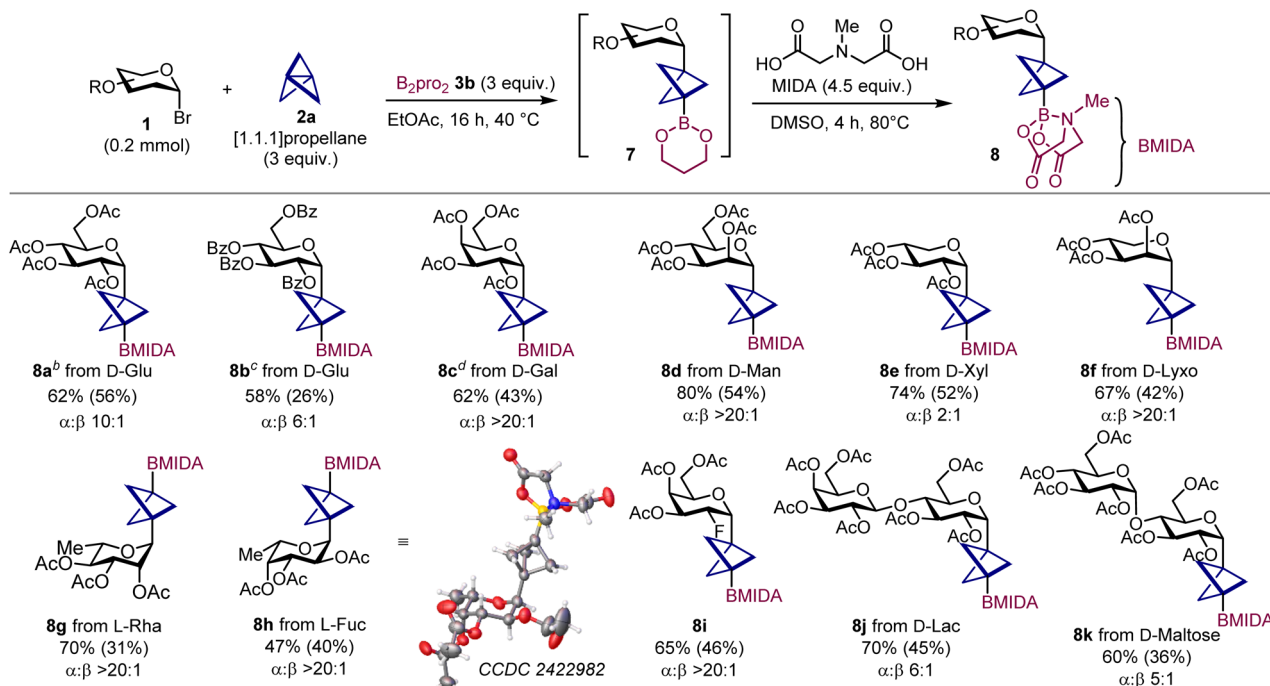
Mechanistic investigation

After having assessed the generality of the developed methods, we aimed to elucidate the involved reaction mechanisms, starting from the hydroglycosylation protocol (Scheme 3a).

To gather insights on a possible thermal initiation pathway, we first studied the behavior of [1.1.1]propellane **2a** upon mixing it with silane ($(\text{Me}_3\text{Si})_3\text{SiH}$) **3a**. Interestingly, we found that these two species can react together to give the addition product **15**, a compound that could also be identified as side-product in the model reaction crude mixture. This addition reaction is completely inhibited in the presence of TEMPO (2,2,6,6-tetramethyl-1-piperidinyloxy) and a TEMPO adduct strongly support the involvement of silyl radicals (for details see SI, Section G).

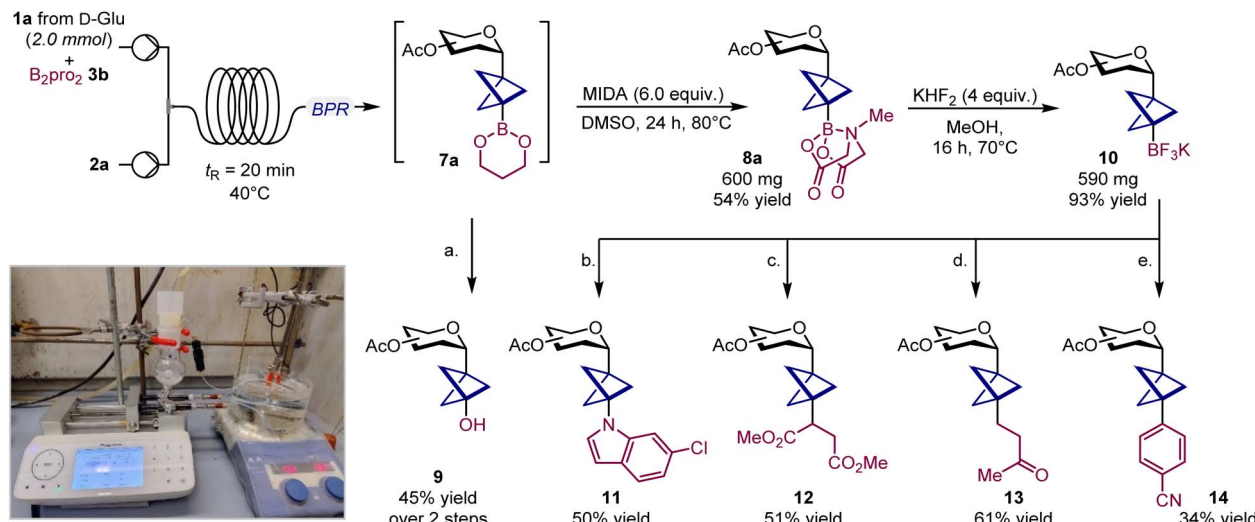
Along this line, when the model reaction was performed in the presence of TEMPO, the formation of the BCP *C*-glycoside **4a** was also completely suppressed, further supporting the radical nature of the transformation. A ^2H -labelling experiment performed using the deuterated silane **3a-d₁** resulted in the formation of glycoside **4a** with 50% deuterium incorporation (as judged by ^1H NMR), which unequivocally demonstrated the role of $(\text{Me}_3\text{Si})_3\text{SiH}$ **3a** as HAT donor. Based on these findings, we propose the following mechanism for the hydroglycosylation reaction (Scheme 3a bottom). A main initiation pathway involves the oxidation of silane **3a** ($E_{\text{ox}} ((\text{Me}_3\text{Si})_3\text{SiH}^+/\text{Me}_3\text{Si})_3\text{SiH} = +0.73 \text{ V vs. SCE in } \text{CH}_3\text{CN})$ ²³ from the photoexcited 4CzIPN* ($E_{\text{red}}^{1/2} (\text{PC}^*/\text{PC}^{\cdot-}) = +1.43 \text{ V vs. SCE in } \text{CH}_3\text{CN}$),^{14b} which upon deprotonation leads to the formation of silyl radical

Table 3 Carboboration of [1.1.1]propellane with glycosyl bromides^a



^a Reactions performed on a 0.2 mmol scale using [1.1.1]propellane **2** as solution in Et₂O (3.0 equiv.), and $B_2\text{pro}_2$ **3b** (3.0 equiv.). Yields outside brackets refer to **7** as determined by ^1H NMR. Yields in brackets refer to isolated products **8**. $\alpha:\beta$ ratios were determined by ^1H NMR analysis of the crude mixture using trichloroethylene as the internal standard. ^b Carboboration run for 2 h. ^c Carboboration run for 5 h. ^d Reaction run using 4 equiv. of both **2a** and $B_2\text{pro}_2$. $B_2\text{pro}_2$: bis(1,3-propanediolato)diboron; MIDA: *N*-methyliminodiacetic acid.





Scheme 2 Reaction scale-up and product derivatization. Scale up performed on 2.0 mmol scale using [1.1.1]propellane **2a** as solution in Et₂O (3.0 equiv.), and B₂pro₂ **3b** (3.0 equiv.). Yields refer to isolated products. (a) Oxone (10 equiv.), K₃PO₄ (3 equiv.), H₂O (5 equiv.), THF, 50 °C. (b) **10** (2 equiv.), 6-chloroindole (1 equiv.), Ir[dF(CF₃)ppy]₂(bpy)PF₆ (2 mol%), Cu(acac)₂ (50 mol%), Cs₂CO₃ (3 equiv.), LEDs λ = 456 nm, 1,4-dioxane. (c) **10** (1 equiv.), Ir[dF(CF₃)ppy]₂(bpy)PF₆ (2 mol%), Na₂HPO₄·2H₂O (3 equiv.), dimethyl fumarate (1.5 equiv.), LEDs λ = 456 nm, THF. **12** was obtained as a mixture with 1 : 1 d.r. (d) **10** (1 equiv.), Ir[dF(CF₃)ppy]₂(bpy)PF₆ (2 mol%), Na₂HPO₄·2H₂O (3 equiv.), Cu(CH₃CN)₄BF₄ (12.5 mol%), methyl vinyl ketone (1.5 equiv.), LEDs λ = 456 nm, THF. (e) Ir[dF(CF₃)ppy]₂(dtbpy)PF₆ (5 mol%), 4-bromobenzonitrile (1 equiv.), **10** (1.5 equiv.), Na₂CO₃ (2 equiv.), Ni(dtbpyp)₂Br₂ (10 mol%), LEDs λ = 440 nm, 1,4-dioxane:DMA, 4 : 1. B₂pro₂: bis(1,3-propanediolato)diboron; BPR: back pressure regulator; MIDA: *N*-methyliminodiacetic acid.

IIIa.²³ Oxidative quenching of the photoexcited 4CzIPN* ($E_{\text{ox}}^{1/2}$ (PC^{•+}/PC*) = -1.18 V vs. SCE in CH₃CN)^{14b} from glycosyl bromide **1** (E_{red} (**1a**/**1a**^{•-}) = -2.45 V vs. SCE in CH₃CN)²⁴ can be excluded on the basis of unfavorable thermodynamics. This mechanistic pathway is supported by Stern–Volmer quenching studies, which revealed (Me₃Si)₃SiH **3a** as the only reaction component able to quench the photoexcited 4CzIPN* (for details see SI, Section G).

Besides, since control experiments revealed that the reaction can also proceed in the absence of light, a further thermal initiation pathway is likely operative. In this regard, we propose that (Me₃Si)₃SiH **3a** reacts with propellane **2** to give silyl radical **IIIa** and BCP radical **IV**. The latter can further react with another molecule of **3a** to give **16** and a second silyl radical **IIIa**. Then, silyl radical **IIIa** can act as a chain carrier undergoing halogen atom transfer (XAT) with glycosyl bromide **1**,²⁵ giving the corresponding glycosyl radical **I** and (Me₃Si)₃SiBr (whose formation was confirmed by GC/MS analysis). Finally, addition of **I** to propellane **2** leads to BCP radical **II**, which subsequently reacts with **3a** to afford the BCP C-glycoside **4** and regenerate silyl radical **IIIa**, thereby sustaining a radical-chain. In this scenario, the side-product **15** is formed through a competitive addition reaction of silyl radical **IIIa** to propellane **2**. Although the reaction can also proceed thermally, the use of 4CzIPN provides an additional pathway to reinitiate the chain reactions once they are terminated, which explains the improved outcome when the reaction is carried out under light and in the presence of the photocatalyst.

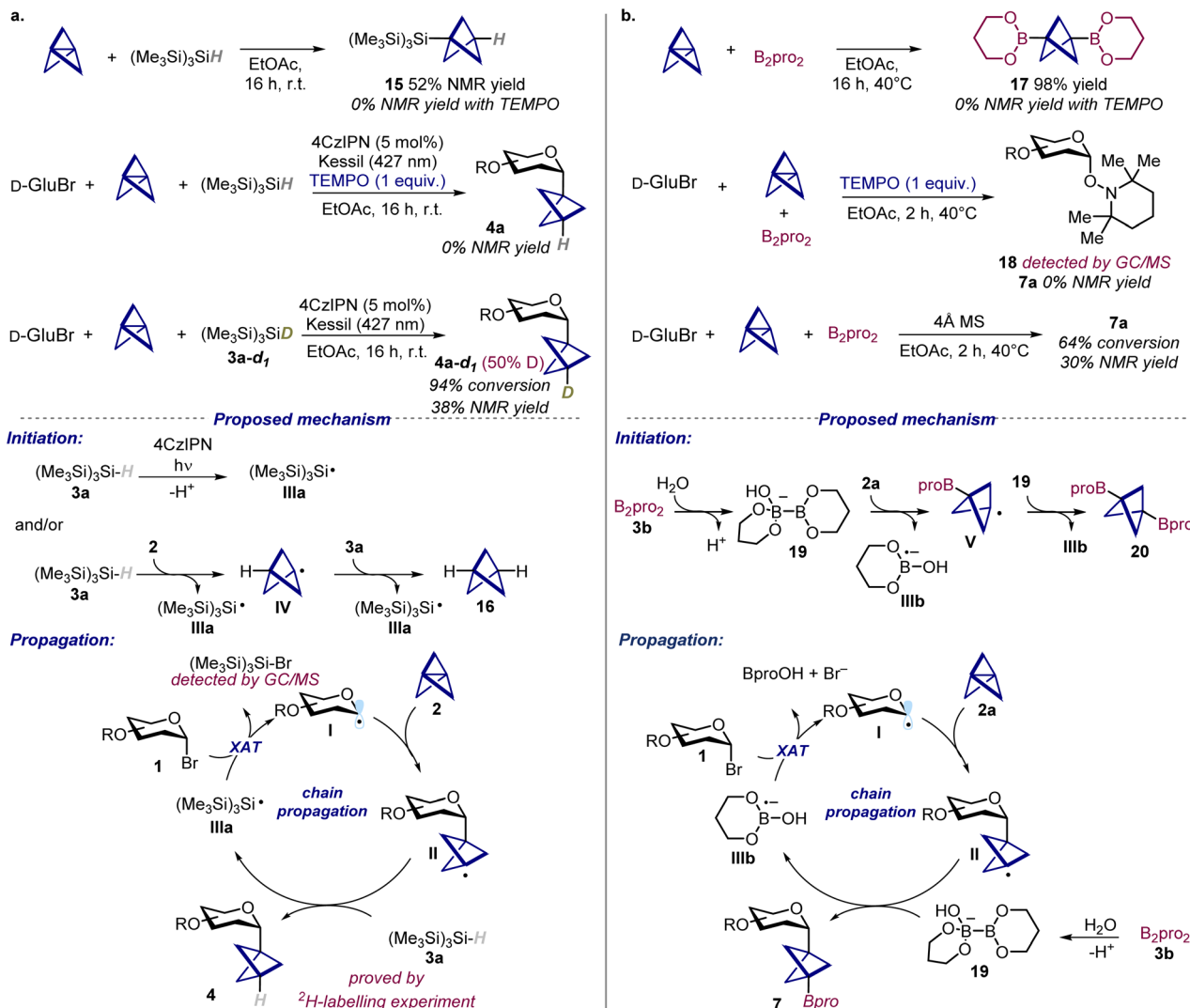
We next investigated the mechanism for the carboboration of propellane (Scheme 3b). Following the same approach, we first demonstrated that the SOMOphile B₂pro₂ can react with propellane **2a** to give the bisboronate **17**, which likely represent

the main initiation pathway for the reaction (*vide infra*). Such addition process, as well as the model carboboration reaction were completely inhibited when performed using TEMPO. In the latter case we were also able to detect by GC/MS the TEMPO adduct **18**, which proves the formation of glycosyl radicals as intermediates. Then, we interrogated the ability of B₂pro₂ to behave as a SOMOphile in the optimized reaction conditions. This diboron(4) compound is able to intercept radical species upon coordination from a Lewis base.^{17b} Here, we showed that the reaction is strongly inhibited when performed using 4 Å molecular sieves, which suggest that water might activate B₂pro₂. Based on these observations, we propose a radical mechanism initiated by reaction between B₂pro₂ **3b** and propellane **2a**. Specifically, we propose that coordination of B₂pro₂ by water might lead to the formation of the active ate-complex **19** that reacts with propellane **2** giving boryl radical anion **IIIb** and BCP radical **V**.^{17b,26} Addition of **V** to **19** then gives the bisboronate side-product **17**.²⁷ On the other hand, we speculate that the resulting boryl radical anion **IIIb** can then undergo XAT with glycosyl bromide **1**, forming glycosyl radical **I**. Finally, subsequent addition of radical **I** to propellane **2** and activated **19** leads to the desired BCP boronate **7** and restores intermediate **IIIb**, thus propagating the chain reaction.

Next, seeking to determine the origin of the α stereoselectivity, we performed DFT calculations investigating the reaction between peracetylated D-glucosyl radical **Ia** and [1.1.1] propellane **2a**. This key reactive intermediate populates three main conformations of the six-membered ring, namely the chair ⁴C₁, the inverted chair ¹C₄, and the boat B_{2,5}.³¹ These conformers exhibit relative free energy values spanning a narrow range of 1.74 kcal mol⁻¹, with the B_{2,5}-**Ia** conformer



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Scheme 3 Mechanistic probes and proposed mechanism for the (a) hydroglycosylation and (b) carboboration of propellanes. 4CzIPN: 1,2,3,5-tetrakis(carbazol-9-yl)-4,6-dicyanobenzene; TEMPO: 2,2,6,6-tetramethyl-1-piperidinyloxy.

being the most stable. They also show similar calculated global electrophilicity ($1.03 < \omega < 1.08$ eV), consistent with nucleophilic α -oxo carbon centered radicals.²⁸ The addition of **Ia** to **2a** to give BCP radical **IIa** is exergonic, with a calculated ΔG^0 of -12.4 and -14.8 kcal mol⁻¹ for the α and β anomer, respectively.²⁹ Experimental data show the α anomer to be the favored one, suggesting that the stereoselectivity is likely determined by kinetic factors. Indeed, among the possible transition states, the lowest energy one was found to be $TS\alpha$, which shows a 4C_1 conformation (incipient C···C bond distance of 2.297 Å, Fig. 1) while favoring α stereoselectivity ($\Delta G^\ddagger = +9.3$ kcal mol⁻¹).³⁰ For comparison, the related $TS\beta$ is associated to $\Delta G^\ddagger = +12.2$ kcal mol⁻¹ for the pathway to β -**IIa** anomer; other TS adopting different conformations (namely 1C_4 or $B_{2,5}$) were found at higher energies for both α and β pathways (SI, section H). Spin density maps of both TS clearly show that the anomeric radical is partially delocalized over the endocyclic oxygen of the sugar ring, an effect that is more pronounced for the $TS\alpha$. This

is indicative of a better orbital overlap between the lone pair of the oxygen atom and the σ^{*} orbital of the incipient C-C bond.¹⁹ Such stabilizing effect – *i.e.* the kinetic anomeric effect in radical glycosylation reactions – accounts for the α selectivity observed in the addition of glucosyl radical to [1.1.1]propellane.

An analogous DFT calculation was performed for the reaction between the peracetylated xylosyl radical **Ie** and [1.1.1]propellane **2a**, which revealed a similar trend of diastereoselectivity with the α -**Ie** anomer being the most favoured by a calculated $\Delta\Delta G^\ddagger$ of 1.5 kcal mol⁻¹. This value is lower than the calculated $\Delta\Delta G^\ddagger$ of 2.9 kcal mol⁻¹ for α -**IIa**, thus being consistent with the lower stereoselectivity experimentally observed for the addition of this xylosyl derivative to [1.1.1]propellane **2a** (see SI, section H for details). These results are significant, as the stereoselectivity of glycosyl radical addition reactions has rarely been investigated by DFT calculations, and our findings align well with the available precedents.³¹

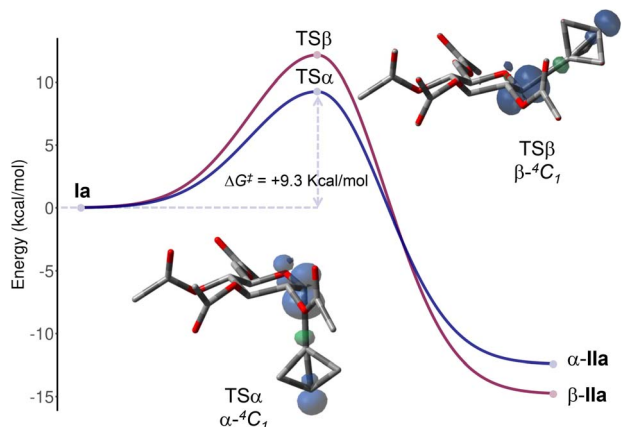


Fig. 1 Energy diagram for the addition of peracetylated glucosyl radical **1a** to [1.1.1]propellane **2**. DFT calculation at M062X/6-311+G**//M062X/6-311+G** level of theory, including a continuum solvation model for Et₂O using the integral equation formalism variant (IEPCM).

Conclusions

In summary, we have developed a mild radical approach for the preparation of saturated analogs of aryl *C*-glycosides. The generality of our strategy was proved in a hydroglycosylation protocol enabling the synthesis of glycosylated BCP derivatives, where the BCP moiety mimics an unsubstituted phenyl ring. By extending this reactivity to the use of diboron(4) SOMOphiles we provided access to difunctionalized glycosylated BCP boronates. This method stands out for its practicality, scalability, and synthetic versatility of the products, which enables straightforward access to a variety of BCP *C*-glycoside derivatives. Experimental and computational mechanistic studies revealed that both protocols proceed through a radical chain mechanism under a Curtin–Hammett scenario, with the stereodetermining addition of glycosyl radicals to [1.1.1]propellane being controlled by the kinetic anomeric effect. Altogether, this study introduces the replacement of planar arenes with highly three-dimensional fragments for the preparation of aryl *C*-glycoside analogues and paves the way for new developments in *C*-glycosylation chemistry.

Author contributions

GG designed the project. GG, AM, SB, PGR performed the experiments. GP performed X-ray analyses. AS performed DFT calculations. LD supervised the work and secured fundings. GG and LD wrote the manuscript with contributions from all the authors. All authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data that support the findings of this study are available free of charge in the supporting information (SI). Supplementary information: includes details of experimental procedures, full

characterization data, copies of NMR spectra, X-ray analysis, and DFT calculations. See DOI: <https://doi.org/10.1039/d5sc07328f>.

CCDC 2422982 contains the supplementary crystallographic data for this paper.³²

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