Oxo-Rhenium-Mediated Allylation of Furanoside Derivatives: A Computational Study on the Mechanism and the Stereoselectivity

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ABSTRACT: Properly substituted tetrahydrofuran (THF) rings are important building blocks in the synthesis of many natural metabolites. Having reliable procedures to control the stereo-selectivity at the THF core while decorating it with different substituents is a fundamental requirement to achieve and fulfill the complexity of nature. We recently reported a new chemical approach to control the stereochemistry in the alkylation and arylation of furanoside derivatives by using a rhenium(V) complex to form an intermediate oxo-carbenium species able to react with proper soft nucleophiles. Here, we describe theoretical calculations, performed at the DFT B3LYP level, to disclose the important mechanistic features which regulate the entire catalytic cycle of the reaction of mono- and disubstituted furanosides with allylttrimethylsilane catalyzed by Re(O)Cl$_3$(OPPh$_3$)(Me$_2$S). Moreover, the key factors governing the allylation step were investigated, confirming that the stereoselectivity, which is independent of the anomeric configuration of starting acetal, mainly arises from the orientation of the substituent at C-4, with only marginal contribution of the substituent at C-5. Finally, puckering Cremer–Pople parameters were used to take trace of the structural modifications throughout the catalytic cycle.

INTRODUCTION

In the realm of natural metabolites, a place of primary importance is held by THF motifs. A combination of the way in which they can be decorated and the control of the stereochemistry on the five-membered ring has served as a formidable challenge for synthetical chemists. To control the chirality on the ring, many different strategies have been elaborated during the years, but one in particular resulted to be the most widely adopted: namely, the venerable Sakurai−Hosomi reaction, which operates on the acetal forms of the tetrahydrofurans by involving strong Lewis acids (e.g., TMSOTf, SnCl$_4$, BF$_3$·Et$_2$O, or TiCl$_4$) to prepare the cyclic five-membered oxo-carbenium ion, and soft, nucleophilic allylsilanes to afford the corresponding allylated THF product. However, the high moisture sensitivity of strong Lewis acids has limited the applicability of this method. This problem was addressed during the years and was overcome by mainly two different strategies reported by Yadav and Friestad. The first one suggested the usage of molecular iodine in CH$_2$Cl$_2$ at cryogenic temperatures, while the second one uses TiCl$_4$ as a Lewis acid again but under milder reaction conditions. Recently, we reported a new chemical opportunity to control the stereochemistry by using a rhenium(V) complex as a Lewis acid additive to obtain a metal-oxo-carbenium complex. To the best of our knowledge, besides the specific analysis for the anisole derivative we reported in our previous paper, no computational studies have been reported regarding the complete catalytic cycle for this reaction.

Seminal work in 2014 from Woerpel and co-workers computationally demonstrated the allylation step on five-membered ring acetals bearing fused rings, showing how subtle structural changes of the fused ring can result in dramatic influences on selectivity. Moreover, in 2021, the computational evidence of glycosyl cations as intermediates in the glycosylation reactions taking place through a S$_N$1-type mechanism has been shown, thus confirming the validity of using this structural representation to describe the reaction.

In the present work, we computationally investigate each step of the catalytic cycle promoted by the rhenium(V) catalyst in the allylation of furanoside derivatives, disclosing the key features which finely regulate the reaction mechanism and, importantly, the selectivity in the allylation step. Moreover, we dedicate special attention to the puckering parameters associated with the conformational changes the structures undergo during the course of the reaction.

COMPUTATIONAL METHODS

The mechanism for the rhenium-catalyzed reaction was investigated using the Gaussian 09 program package in the framework of the
density functional theory (DFT). All the calculations were carried out using, as a hybrid functional for DFT calculation, the Becke three-parameter hybrid exchange functional (B3) in its variation provided by the Lee, Yang, and Parr correlation functional (LYP). The basis set was specifically selected for each atom, considering the type and neat differences in the electronic distribution. Specifically, H, C, O, and F atoms were described by using the 6-311+G(d,p) basis set, whereas P, S, and Cl atoms were described using the 6-311+G(2df,p) basis set. Differently, the LanL2DZ effective core potential basis set was selected for the Re and Cu atoms. Open-shell calculations using the UB3LYP functional were performed when Cu(II) species were involved, to fulfill their doublet spin-state nature. Further calculations on benchmarking of theory can be found in the Supporting Information. For each structure, frequency calculations were performed to confirm the effective minimum or transition-state (TS) nature of the optimized structure. Moreover, natural population analysis was performed, considering the charge stabilization for the oxygen (O-1) and the acetal carbon (C-2) atoms during the formation of the oxo-carbenium cation and the allylation process. The key steps of the mechanism are reported in Scheme 1, together with the important features we highlighted during the calculations. The analysis of the mechanism will be divided into four main sections that retrace the key steps of the reaction: the coordination of the THF acetal by the rhenium active species, the formation of the oxo-carbenium ion, its allylation, and the regeneration of the catalyst.

We also dedicated a special effort to model the formation of the catalytically active Re(V) species from the rhenium catalyst. Indeed, the reaction initiates with the Cu(OTf)2 displacement of Me2S from the rhenium complex Re(O)Cl3(OPPh3)(Me2S)A. This reaction step was investigated computationally, showing an activation energy for the ligand exchange of 11.2 kcal mol−1 from the adduct between the two reagents A + Cu(OTf)2 Adduct, with an increase of stability for the obtained B product of nearly 12 kcal mol−1 from the starting materials (Chart 1). The driving force for this reaction step is due to the higher thiophilicity of Cu compared with that of Re.

**Coordination of THF-Methoxy Acetal 1 by the Reactive Re Species B.** The catalytically active 12e− Re-complex B thus formed reacted with the 2-methoxymethoxytrihydrofuran 1 to produce a THF−Re complex 2 via a low-barrier (6.9 kcal mol−1) TS, TS-1 (Chart 1). During this step, an almost unchanged charge on the C-2 carbon of the THF ring (from +0.429 to +0.436) has been observed; the same situation occurs also on the O-1 oxygen, which showed an unaltered value of charge (from −0.626 to −0.619) but of the opposite sign (Scheme 1). In TS-1, substrate 1 directs only the aglycone oxygen atom of the −OMe residue toward the Re metal center by maintaining the more stable exo-anomeric orientation for the methyl group during the formation of the interaction (3.09 Å). The THF−
oxygen remains far from the Re coordination sphere; in fact, when we considered a possible chelation of the two oxygen atoms of \( \text{Rn} \) to \( \text{Re} \), less stable structures were found both for the complex \( \text{TS-1} \) and \( \text{TS-2} \). This particular behavior was observed for all further steps of the catalytic cycle and thus has been excluded as a viable pathway.

The \( \text{Re} \)–\( \text{OMe} \) distance registered during the TS (3.09 Å) was significantly shortened to 2.37 Å in the intermediate \( \text{TS-1} \). Moreover, the evaluation of puckering parameters showed no considerable modification in conformation by moving from \( \text{TS-1} \) to \( \text{TS-2} \) and finally to \( \text{TS-3} \); the \( \text{E} \) envelope conformation remains unchanged during all these steps (see the Puckering polar plot in the Supporting Information for more details), as well as the C-2/OMe distance only slightly lengthened from 1.41 Å in \( \text{TS-1} \) and 1.48 Å in \( \text{TS-2} \). This indicates that no modification in the structure of the THF acetal is necessary during the direct coordination of the \( \text{-OMe} \) substituent to the rhenium metal center (see Table S2 in the Supporting Information for more details). Another important detail that arises from these calculations is related to the pyramidalization of the C-2 atom, which indirectly indicates the hybridization at C-2. To describe this property, we decided to look for the sum of the angles between the substituents at C-2 not involved in the reaction (namely, O-1, H, and C-3). By measuring the angle between them and summing the results, we can conclude that if the sum is close to 360°, the hybridization at C-2 is almost sp\(^2\); if the sum is close to 330°, the hybridization at C-2 is almost sp\(^3\). The results we got go from 327.0° in \( \text{TS-1} \), to 328.1° in \( \text{TS-2} \), and then to 329.5° in \( \text{TS-3} \), thus indicating that TS-1 has the only role in coordinating \( \text{Rn} \) to the Re-complex with no change in the hybridization at C-2, with the angle values being compatible with sp\(^2\) hybridization of C-2 (see Table 1).

**Formation of Oxo-Carbenium Ion 3.** Complex \( \text{TS-2} \) then moves toward the rate-determining TS-2 with an energy barrier of 16.1 kcal mol\(^{-1}\). \( \text{TS-2} \) features an increased positive charge on C-2, from +0.448 in \( \text{TS-1} \) to +0.541, and a significantly less negative charge on O-1, from -0.598 to -0.401. The complete breaking of the crucial MeO–THF bond, increasing in distance from 1.48 to 2.98 Å, and the concomitant shortening of the MeO–Re distance to 1.99 Å indicate a product-like nature of TS-2. Once the energy barrier has been overcome, two charged species are formed, namely, the cationic oxo-carbenium 3 (charges of +0.559 and -0.369 for C-2 and O-1, respectively) and the anionic rhenium complex C. This step also highlights the S\(_\text{N}1\) mechanism of this reaction: the slow ionization to form the oxo-carbenium ion is achieved in TS-2 during the rate-determining step, and only in a second time, the fast reaction between the intermediate cation and the nucleophile occurs (see Chart 3).

During this step, the THF ring reverts the envelope conformation from \( \text{E} \) in \( \text{TS-2} \) to \( \text{E} \) in 3, passing through an intermediate conformation.

**Table 1. Pyramidalization Angles Related to the Hybridization at the C-2 Atom for the Catalytic Cycle**

|       | Sum of angles of pyramidalization\(^a\) (°) | Hybridization at C-2 |
|-------|-----------------------------------------|---------------------|
| 1     | 327.0                                  | sp\(^3\)            |
| TS-1  | 328.1                                  |                     |
| 2     | 329.5                                  | sp\(^2\)            |
| TS-2  | 360.0                                  |                     |
| 3     | 360.0                                  |                     |

\(^a\)Sum of all angles on C-2 \( \text{(O1-C2-H + O1-C2-C3 + C3-C2-H)} \).
Chart 2. Electronic Energy Profile with ZPE Correction (kcal/mol) for the Regeneration of Re Catalyst Species B$^\text{a}$

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B3LYP/6-311+G(d,p)&6-311+G(2df,p)&LanL2DZ/CH$_2$Cl$_2$(PCM)//B3LYP/6-311+G(d,p)&6-311+G(2df,p)&LanL2DZ.
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Chart 3. Electronic Energy Profile with ZPE Correction (kcal/mol) for the Allylation of Oxo-Carbenium ion 3$^\text{a}$

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In blue are reported the conformational descriptors for the five-membered ring. CH$_2$Cl$_2$(PCM)-B3LYP/6-311+G(d,p)&6-311+G(2df,p).
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The pyramidalization of the C-2 atom which depends on the hybridization of C-2: the sum of angles moves from 329.5° to 2 in 360.0° in TS-2, a compatible value with the sp² hybridization of C-2 in 3 (see Table 1).

Further interaction of 3 with allytrimethylsilane afforded the corresponding allylated THF. The trimethylsilyl cation, which originated from the allylation step, combines with anionic complex C during the neutral TS, TS-3 (Chart 2). The incoming cation directs toward the oxygen atom of the coordinated MeO-residue, with the assistance of the nearby coordinated chlorine atom. This transformation shows an energy barrier of 10.9 kcal mol⁻¹ (TS-3) with the silicon atom shared between chlorine and oxygen atoms (bond lengths of 2.44 and 2.04 Å, respectively). The TS-3 evolves in the next Re-complex D, with a completely formed Si–OMe bond (1.74 Å) and a stabilization of nearly 23 kcal mol⁻¹ from the previous TS (Chart 2). Decomplexation of the MeO–TMS ether from the Re-complex D restores the 12e Re-complex B, which re-enters the catalytic cycle. This last step occurs through the TS-4 TS, with an activation energy of only 3.5 kcal mol⁻¹.

The selectivity appears to be strongly dependent on steric hindrance provided by the silylgroup at C-5. In order to minimize the steric interaction with the pentacyclic ring, the TMS group of the other substrate at C-5 is oriented outward. Thus, both disubstituted faces of ion 5 are accessible, but the shielding of the upper face due to the inward OTMS group orientation results in a calculated energy difference between the two most populated TSs of 0.8 kcal mol⁻¹, corresponding at 22 °C to a 74.3:25.7 trans/cis-product ratio. This selectivity is in agreement with the experimental 60:40 value observed in the allylation of compound 5, as reported by Woerpel and co-workers. The distance between the oxo-carbenium C-2 carbon and the terminal allylic one resulted to be nearly the same in both the approaches (~2.2 Å). Moreover, the allytrimethylsilane distorts the structure of the oxo-carbenium ion in a different way, according to the direction of its approach: the trans-approach favors the 3°, 7° conformer TS leading to the cis-product. Finally, the best conformation of the two possible attacks on the oxo-carbenium ion will be represented in a tridimensional structure in order to highlight the important features which regulate the observed selectivity.

Table 2. Pyramidalization Angles Related to the Hybridization at the C-2 Atom for the Allylation Step

| Sum of angles of pyramidalization(°) | Hybridization at C-2 |
|--------------------------------------|----------------------|
| 3 + allyl-TMS Adduct | 359.0 | sp² |
| TS-S | 352.8 | sp² / sp³ |
| 4 + TMS* | 326.9 | sp³ |

“Sum of all angles on C-2 (O1–C2–H + O1–C2–C3 + C2–C3–H).”

previous paper³ or taken from the literature.¹⁹,²⁰ Specifically, we considered different situations depending on the substituents at C-4 and C-5 of the oxo-carbenium ion: we started with the mono-substituted glycosyl cations 5 and 6 to then increase the complexity of the system to disubstituted intermediates 7 and 8 (Figure 1). However, the energy change without losing steric information, we decided to simplify the TBS or TBDPS moieties with a TMS group. Moreover, for species 5, 7, and 8, since there is the possibility of different orientations of the substituent at the C-5 position, we evaluated all the three staggered rotamers by reporting in the tables each of the minima as T1 (gt-rotamer), T2 (gt-rotamer), and T3 (gt-rotamer) (see Figure S1 in the Supporting Information for more details). Finally, the best conformation of each of the two possible attacks on the oxo-carbenium ion will be represented in a tridimensional structure in order to highlight the important features which regulate the observed selectivity.

According to the published experimental results,³,¹⁴ the stereochemistry of the anomeric center of the furanoside which originates the oxo-carbenium species is irrelevant to the observed selectivity; thus, only the allylation step determines the selectivity. In fact, it has been demonstrated by Woerpel and co-workers that the allylation step along the reaction pathway is governed by the Curtin–Hammett principle.²¹ Hence, for these substituted substrates, only the corresponding stereoisomeric TSs were located and the theoretical diastereomeric product ratio was determined by the relative energies of the two TSs (i.e., a and b) originating from the common oxo-carbenium intermediate (Figure 1).

For the C-5 mono-substituted oxo-carbenium ion 5, DFT calculations identified three TSs leading to the trans,2,5-disubstituted product and three TSs leading to the cis-product. The most stable TS of each approach (Figure 2) shows the oxygen of the side chain above the ring and close to the oxonium center to allow a stabilizing interaction with the oxo-carbenium moiety (T1 conformation). The selectivity appears to be strongly dependent on steric hindrance provided by the silylgroup at C-5. In order to minimize the steric interaction with the pentacyclic ring, the TMS group of the other substituent at C-5 is oriented outward. Thus, both disubstituted faces of ion 5 are accessible, but the shielding of the upper face due to the inward OTMS group orientation results in a calculated energy difference between the two most populated TSs of 0.8 kcal mol⁻¹, corresponding at 22 °C to a 74.3:25.7 trans/cis-product ratio. This selectivity is in agreement with the experimental 60:40 value observed in the allylation of compound 5, as reported by Woerpel and co-workers. The distance between the oxo-carbenium C-2 carbon and the terminal allylic one resulted to be nearly the same in both the approaches (~2.2 Å). Moreover, the allytrimethylsilane distorts the structure of the oxo-carbenium ion in a different way, according to the direction of its approach: the trans-approach favors the E=1/1, 1° puckering, while the cis-one prefers the opposite 1°, 1° conformer (see Table S4 in the Supporting Information for more details).

Regarding the C-4 mono-substituted oxo-carbenium ion 6, we do not have the three possible orientations reported for the case above since there is no methylene functionality that can result in an easier rotation of the dihedral angle. For this reason, we analyzed only one conformer for both the cis- and trans-approach. In both the situations, we observed that the silylgroup at C-4 is oriented outward to
reduce the steric hindrance. Moreover, in agreement with Woerpel’s observations, in both the TSs, the C-4 silyl ether group is pseudoaxially oriented (Figure 3). When the conformations with a pseudoaxial substituent at C-4 were investigated, they were found not to be the local minima on the potential energy surface by converging to the pseudoaxial structures during the optimizations. The calculated energy difference between the two TSs is 1.9 kcal/mol, corresponding at 23 °C to a 95.9:4.1 cis/trans product ratio. This result is in good agreement with the experimental value of 94:6 determined for the allylation of 6, as reported by Woerpel and co-workers.

In this new case, the distance between the oxo-carbenium C-2 carbon and the terminal allylic one resulted to be slightly longer than the previous one, yet is nearly identical in both approaches (∼2.3 Å). Also, in this case, the approach of the allyltrimethylsilane distorts the structure of the oxo-carbenium but not in an opposite manner for the two directions of the approach like in the previous situation. The trans-approach favors the E$_5$/T$_5$ puckering, while the cis-one prefers the $^4T_3$ like in the previous system (see Table S5 in the Supporting Information for more details). This variation is mainly due to the different location of the substituent on the oxo-carbenium structure, together with the higher conformational rigidity imparted by the pseudoaxially oriented silyl ether group at C-4.

The computational and experimental results on the allylation of the above reported oxo-carbenium ion intermediates clearly indicate the ability of the substituent at C-5 to promote the trans-allylation, whereas the alkoxy substituent at C-4 is significantly cis-orienting. These abilities may result in a match or mismatch when the 4,5-disubstituted oxo-carbenium ions 7 and 8 are considered.
In particular, a mismatch can be predicted in the case of the cis-4,5-disubstituted oxo-carbenium ion 7. The DFT calculations predicted the all-cis trisubstituted THF product as the preferred diastereoisomer but with a lower selectivity than in the case of 6. Indeed, the calculated cis/trans ratio (considering all possible orientations on the C-5 silyl ether) is 77.3:22.7, in good agreement with the available experimental data from our reactions, which show a diastereomeric ratio of 73:27.

The most stable TSs for the cis and trans allylation of oxo-carbenium ion 7 (shown in Figure 4) differ by 0.6 kcal/mol at −30 °C. The most stable TSs for the cis and trans allylation of oxo-carbenium ion 7 (shown in Figure 4) differ by 0.6 kcal/mol at −30 °C. In both the approaches, the C-4 silyl ether group is pseudoaxially oriented as in 6, but the effect it exerts is directed toward the side chain at C-5. Indeed, its vicinal presence forces the C-5 silyl ether in the pseudoequatorial orientation, outside the heterocyclic ring, favoring the T3 conformation and thus reducing the contrast in the approach of nucleophile toward the cis-allylation. Vice versa, the trans-approach prefers a T2 conformation, given the impossibility of orienting the oxygen of the side chain at C-5 above the ring and close to the oxonium center like in oxo-carbenium 5. The pseudoequatorial orientation remains the most favored also for the trans-approach. Since the orientations of the substituents are strongly
different with respect to the previous mono-substituted cases, we should expect a strong variation in terms of puckering parameters. This is particularly evidenced in the trans-approach, where the $T_1$ conformation assumes the $3_T$ puckering as for $5$, but the other most populated one (namely, $T_2$) shows the $4_T$ puckering, like in the previous trans-allylation in $6$. Regarding the cis-approach, the $4_T$ puckering remains again the most favored (see Table S6 in the Supporting Information for more details). It thus seems that the strong role in controlling the puckering derives from the orientation of the substituent in C-4, rather than the one in C-5, in accordance with the 1,3-selectivity (2,4-one with our numbering system) described by Woerpel and co-workers.\(^{19}\) This observation can also be understood by considering a stabilizing H-bond interaction between the polarized terminal vinylic proton and the oxygen atom at C-4, which is present only if the approach is on the same side of the substituent at C-4.

As a final observation, the distance between the oxo-carbenium C-2 carbon and the terminal allylic one resulted to be similar to the ones reported for $6$ ($\sim 2.3$ Å). Differently from oxo-carbenium $7$, the opposite orientation of the substituent at C-4 in $8$ should reinforce the 1,3-orienting effect (2,4-in our numbering system) in the allylation of the trans-4,5-disubstituted oxo-carbenium ion. According to our experimental results and calculations, synergistic cooperation of the C-4 and C-5 trans-substituents allows almost complete diastereoselectivity. The calculated energy difference between the two most populated TSs is 3.6 kcal/mol, corresponding to a 99.9:0.1 ratio at $-30^\circ$C between the trans- and cis-products. This result is in good agreement with the experimental value of 99:1 determined for the allylation of $8$ as reported in our recent publication.\(^{7}\) The calculated most stable TSs for the cis and trans products are shown in Figure 5. Similar to the last case, here also, we can analyze the orientation of the substituents already installed on the oxo-carbenium ring in detail. In the trans-approach, the $T_1$ conformation remains the most stable one, with both the substituents oriented in pseudoequatorial positions. The puckering remains $3_T$, the classically favored one for the trans-approach that we observed also in the above oxo-carbenium $5$ (see Table S7 in the Supporting Information for more details). Regarding the cis-approach, $3_T$ puckering is assumed only in the most preferred TS conformer (i.e., $T_3$) by showing now a general behavior in bringing the substituents in a bis-pseudo equatorial position. However, the bis-pseudoequatorial one remains still possible in the other less populated conformers (i.e., $T_1$ and $T_3$), where $5_T$ puckering is now preferred (see Table S7 in the Supporting Information for more details). Nevertheless, the stabilization provided by the interaction between the oxo-carbenium moiety and the oxygen of the side chain above the ring is not still strong enough to revert the selectivity like in the first case we analyzed with oxo-carbenium $5$. Moreover, also in this case, the stabilizing H-bond interaction between the polarized terminal vinylic proton and the oxygen atom at C-4 appears favorably, which now directs toward the trans-approach. Finally, the distance between the anomeric carbon and the terminal allylic one resulted to be in all cases similar to the previously reported one ($\sim 2.3$ Å).

### CONCLUSIONS

In conclusion, DFT computational studies were able to assist in the unveiling of the energetic profile for the rhenium-catalyzed allylation reaction of five-membered ring acetals. Importantly, critical steps of the reaction including the rate-determining step in the ionization toward the formation of oxo-carbenium were revealed. Moreover, all the TSs of the diastereoisomeric allylation step were located by showing the key factors which govern the stereoselectivity of the reaction independently from the anomeric configuration of the starting acetal. The theoretical data supports the experimental results reported by Woerpel and co-workers and is proven also in our...
recent publication. In particular, we confirmed the conclusions advanced by Woerpel, where the diastereoselection in C-glycosylation reaction of ribose-derived analogues mainly arises from the relative orientation of the substituent at C-4, given the only marginal, although not negligible, the contribution of the one at C-5.

## EXPERIMENTAL SECTION

### Computational Methods.

For the catalytic cycle, all the structures were optimized using the Gaussian 09 program package, using the B3LYP functional and a differentiation basis set according to the atom type. Specifically, 6-311+G(d,p) was used for H, C, O, and F, and 6-311+(2d,2p) for P, S, and Cl. The effective core potential basis set LanL2DZ was used for Re and Cu atoms. Calculations performed in presence of Cu species were run using UB3LYP, to fulfill the doublet nature of the spin state for Cu(II) species. All the optimizations were performed in vacuo. Frequency calculations were performed for each species in order to confirm the effective minimum or TS nature of the optimized structures. Moreover, IRC calculations were also used to confirm that all TS structures are linked with that of the reactant, intermediate, or products. With the optimized geometries, single-point energy calculations in dichloromethane were performed using the PCM and the same basis set used in the optimization step. All data reported for the catalytic cycle are referred to this level of theory. Moreover, Cremer–Pople conformational analysis was performed in order to identify the puckering parameters which describe the tetrahydrofuran ring as a result of the structural stability. Regarding the computational investigation of the diastereoselectivity in the alkylation step, we adopted a similar level of theory. We located and optimized the TSs in the reaction step with allytrimethylsilane by always using the B3LYP functional and a different basis set: 6-311+G(d,p) was used for H, C, and O atoms, while 6-311+G(2df,p) was used to account for the Si atom. Tight convergence criteria for the SCF cycles have been adopted (SCF = tight). Moreover, the effect of the solvent dichloromethane was directly considered during the optimizations by means of the PCM. Frequency calculations to extract thermochromical parameters and corrections were performed at the specific reaction temperature reported in the papers, and the theoretical product distributions were obtained by applying Boltzmann population analysis. Conformational analysis for the gauche orientations of the substituent around the CH2-chain at the C-5 position was manually performed by identifying three possible relative minima as conformers T1, T2, and T3. Finally, we decided to simplify the TBS or TBDPS moieties with a computationally cheaper TMS group.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.joc.2c00393.

Benchmark with different levels of theory; conformational and structural features of the substituted oxocarbenium species; tables with energies, distances, and puckering parameters for the alkylation of the oxonium species; puckering polar plot; and summary of energies for the optimized structures (PDF)

Cartesian coordinates of the structures, together with the value for the electronic energy and frequencies for the optimized species (XYZ)

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