Original Article

Stereoelectronic Properties of 1,2,4-Triazole-Derived N-heterocyclic Carbenes - A Theoretical Study

Nguyen Van Ha*, Doan Thanh Dat, Trieu Thi Nguyet

Faculty of Chemistry, VNU University of Science, 19 Le Thanh Tong, Hanoi, Vietnam

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Abstract: A theoretical study on stereo and electronic properties of a series of six 1,2,4-triazole-derived carbenes bearing different N4-substituents, namely isopropyl (1), benzyl (2), phenyl (3), mesityl (4), 2,6-diisopropylphenyl (5) and 1-naphthyl (6), has been carried out. Structures of the six carbenes were first optimized using Gaussian® 16 at B3LYP level. Their molecular geometries and electronic structures of the frontier orbitals were examined. The results suggest the similarity in nature of their HOMOs, which all posses σ symmetry with respect to the heterocycle and essentially be the lone electron pair on the C_carbene. Steric properties of the NHCs was also quantified using percent volume buried (% Vbur) approach. The NHC 1 with isopropyl N4-substituent was the least bulky one with % Vbur of 27.7 and the most sterically demanding carbene is 6, which has large 2,6-diisopropylphenyl substituent (% Vbur = 38.4). Interestingly, the NHCs with phenyl and 1-naphthyl N4-substituents display flexible steric hindrance due to possible rotation of the phenyl or 1-naphthyl around the N-C single bond. Beside stereoelectronic properties of the NHC, topographic steric map of their complexes with metal were also investigated.

Keywords: N-heterocyclic carbene, triazolin-5-ylidene, stereoelectronic properties, percent volume buried.

1. Introduction

In the past few decades, N-heterocyclic carbene (NHC) has remarkably transformed from a merely curiosity-driven laboratory discovery into an essential class of ligand in organometallic chemistry[1-6]. NHC popularity and widespread applications can be attributed to their excellent turnability of steric and electronic properties by changing their N-substituents or the carbene backbone itself [7-8].

Among the four classical type of N-heterocyclic carbenes (Figure 1), imidazole and 1,2,4-triazole derived carbenes are isoelobal since the latter is formed by a mere substitution of a CH group in the earlier by a nitrogen atom. Despite their similarity in electronic and steric
properties, it is surprising to notice that the chemistry of 1,2,4-triazole derived carbene is much less explored[9-14] compared to their imidazole-derived cousin.

![Generic structures of the 4 type of classical N-heterocyclic carbenes derived from imidazole (a), benzimidazole (b), 1,2,4-triazole (c) and imidazoline (d).](image)

As a part of our ongoing effort to explore the chemistry and potential application of 1,2,4-triazole derived carbene [15,16], we present in this work the theoretical study of steric and electronic properties of a series of six 1,2,4-triazole derived carbene bearing different N-substituents (Figure 2).

![Structures of the NHCs in this work.](image)

This theoretical study is expected to provide understanding on the stereoelectronic of the triazole-derived carbene under investigation, and hence provide guidance to the choice of N-substituents for the follow-up experimental work on the design of triazole-derived carbene complexes for catalysis application and drug development.

2. Methodology

The All the carbenes under studied were first optimized using Gaussian® 16 at B3LYP level [17-20]. The 6-31G(d) basis set were employed for all atoms [21,22]. The nature of the stationary optimized points was confirmed to represent minima on energy potential surface by frequency analysis. Kohn-Sham orbitals were obtained directly from these calculations.

The steric hindrance of carbenes and their topographic steric maps are analysed using the web tool SambVca 2 developed by Luigi Cavallo [23]. The optimized structures were taken as input for the calculations. Occupation of the coordination sphere by the carbene, percent volume buried $V_{\text{bur}}$, was calculated using a ghost metal atom coordinated by the carbene with metal-carbon distance of 2.01Å. Topographic steric maps of the carbene on their metal complexes were generated using the same SambVca 2 tool.

3. Results and discussion

3.1. Geometry of the carbenes

Singlet-state gas-phased optimized geometries of 1–6 are shown in Figure 1. Selected bond lengths and bond angles are listed in Table 1.

All the singlet states of all the 6 carbenes are in perfectly planar geometries. The N1–C5 and N4–C5 (N–C_{carbene} bonds) are in the 1.348–1.353Å and 1.378–1.386Å ranges. The $\angle$N1C5N4 angles at the carbene ranging from 100.0–100.2°.

Noted that, since the chemistry of 1,2,4-triazole derived carbene remains almost unexplored, hence there exist very few experimental parameters for this type of compounds. One example is the first know stable triazole-derived carbene, 1,3,4-triphenyl-1,2,4-triazol-5-ylide [24]. In its molecular structure, the N–C_{carbene} distances are 1.351(3)Å and 1.373(4)Å for N1–C5 and N4–C5, respectively. The N_{carbene}N angle ($\angle$N1C5N4) of 100.6(2)°.
was observed. In general, the theoretically calculated parameters are closely resemble the reported experimental values for 1,3,4-triphenyl-1,2,4-triazol-5-ylide.

**Figure 3.** Optimized geometries of 1-6.

**Table 1.** Selected bond lengths (Å) and angles (°)

| Parameter | 1   | 2   | 3   | 4   | 5   | 6   |
|-----------|-----|-----|-----|-----|-----|-----|
| N1-C5     | 1.353 | 1.353 | 1.348 | 1.352 | 1.350 | 1.350 |
| N4-C5     | 1.378 | 1.379 | 1.386 | 1.384 | 1.383 | 1.386 |
| N1-N2     | 1.385 | 1.387 | 1.388 | 1.389 | 1.388 | 1.388 |
| N2-C3     | 1.302 | 1.301 | 1.299 | 1.300 | 1.299 | 1.299 |
| C3-N4     | 1.378 | 1.377 | 1.383 | 1.382 | 1.380 | 1.384 |
| N1C-CN4   | 100.2 | 100.1 | 100.2 | 100.0 | 100.2 | 100.0 |

It is interesting to notice the differences in the orientation of the aromatic N-substituent plane with respect to the plane of the triazole heterocycle ring. The mesityl (in 4) and 2,6-diisopropylphenyl (in 5), due to the steric bulk of the methyl and isopropyl substituents, are nearly perpendicular to the triazole ring, forming dihedral angles of 77° and 90°, in 4 and 5 respectively. On the other hand, the phenyl in 3 and 1-naphthyl group (in 6) are more flexible and can rotate along the N-C-aromatic bond. This is evidenced by dihedral angles of 26° and 51° observed in 3 and 6, respectively. The influence of flexible orientation of the aromatic substituents with respect to the triazole on stereoelectronic properties of the respective carbene will be worth examining more details (*vide infra*).

**Figure 4.** Flexible orientation of phenyl and 1-naphthyl ring with respect to the heterocycle.

### 3.2. Electronic properties of the carbenes

Surfaces of the highest energy occupied molecular orbital (HOMO) and lowest energy
unoccupied molecular orbital (LUMO) of the carbenes are shown in Figure 5 and their energy levels are plotted in Figure 6.

It can be noted that the LUMOs have the shape and spacial extension varies from one carbene to another. Overall, the LUMOs are largely localized on the aromatic ring of the substituents. There is little to none contribution from the triazole ring to the carbenes LUMOs. In fact, the lowest \( \pi \)-unoccupied orbital of the triazole fragment, which in nature are essentially \( p \) orbital of the \( C_{\text{carbene}} \), lie relatively higher in energy. Such orbitals are indeed LUMO+3 for 1, LUMO+5 for 2, 3, 4, 5 and LUMO+6 for 6.

![Figure 5. Shape of frontier (HOMO and LUMO) molecular orbitals for 1-6.](image)

![Figure 6. Energy level of HOMO (blue) and LUMO (red) orbitals of the carbenes.](image)

In contrast the the LUMO, the HOMO orbitals are mainly localized on the triazole ring for 1, 2, 3, 4 and 5. In case of 6, small contribution from the naphthyl ring was spotted (Figure 5). All the HOMO orbitals has \( \sigma \) symmetry with respect to the NHC plane and corresponds to the lone pair of the carbene carbon atom. The high energy nature of the lone pair of \( C_{\text{carbene}} \) atom suggests that these orbitals would involve in formation of bonding between carbenes and transition metal ions.

Close examination of the frontier orbitals reveals a significant difference in LUMO energy level, which is in line with the vast difference in their nature and spatial extension. On the other hand, the energy of HOMO energy levels only slightly varies from a NHC to another, and lie in the range from -5.83 to -6.04eV. The HOMO of 1 is highest in energy \( (E_{\text{HOMO}} = -5.83\text{eV}) \) due to the destabilization by positive inductive effect (+I) from the electron donating isopropyl group. Their HOMO energy similarity suggests a relatively similar donor strength for these carbenes.
3.3. Steric properties of the carbenes

Steric hindrance generally play a dominant role in defining metal complex reactivities, especially in catalysis. There exist several methodologies to evaluate ligand steric hindrance, such as Tolman cone angle [25,26], solid angle measure [27], angular symmetric deformation coordinate [28], ligand repulsive energy parameter [29] and percent volume buried [23,30]. Among these, percent volume buried are a modern approach, which is convenient to use well accepted in the organometallic research community. Percent volume buried (%Vbur) of a ligand is defined as the percentage of the metal center coordination sphere occupied by that particular ligand (Figure 7).

Figure 7. Ligand occupation of the coordination sphere, principle of %Vbur calculation.

Input for calculation of %Vbur of a ligand is solid-state X-ray determined molecular structure of its metal complexes or the optimized geometries of the ligand with a ghost metal ion placed at a certain distance from the C-carbene. Calculation can be performed using SambVca 2, a web-based tool by Luigi Cavallo.

%Vbur values for the six NHCs are listed in Table 2. It can be noted that %Vbur is heavily depend on the nature of the N-substituents. The less bulky isopropyl group form a relatively compact carbene 1 with %Vbur of 27.7. Slightly higher %Vbur values are found for NHC with benzyl (2, 29.7), 1-naphthyl (6, 29.7), mesityl (4, 31.1) and phenyl (3, 31.3) N-substituents. In line with chemistry intuition, bulky 2,6-diisopropylphenyl group gives rise to the carbene 6, which possesses the highest steric hindrance with %Vbur of 38.4.

Table 2. Percent volume burried (%Vbur) of the NHCs

| NHC | %Vbur | NHC | %Vbur |
|-----|-------|-----|-------|
| 1   | 27.7  | 4   | 31.1  |
| 2   | 29.7  | 5   | 38.4  |
| 3   | 31.3  | 6   | 29.7  |

It has been pointed out that the phenyl (in 3) and naphthyl (in 6) substituents can rotate around the N-C single bond, and hence possess a certain degree of flexibility in term of their relative orientation to the triazole ring. Such rotational flexibility is expected to translate into a flexible steric bulk for 3 and 6. In order to probe that sterical flexibility in more details, %Vbur of 3 and 6 were calculated using hypothetical structures formed by rotating the respective substituent along the N-C bond. The %Vbur of the ligand is then plotted against the dihedral angle between the triazole ring and plane of the aromatic substituent (Figure 8).

Figure 8. Changing of %Vbur for 3 and 6 as the phenyl and naphthyl substituent rotates around the N-C bond.

As shown in Figure 8, the %Vbur for both 3 and 6 vary as the phenyl and naphthyl plane rotate around the C-N single bond. For carbene 3, a minimum steric bulk %Vbur of 28.5 is achieved with the phenyl lies perpendicular to
the triazole ring (θ = 0°). As the phenyl ring rotate, %V_{bur} for 3 gradually increases and reaches the maximum of 31.8 when the two planes are coplanar (θ = 0° or 180°). On the other hand, due to unsymmetrical nature of the naphthyl substituent, as it rotates around the C-N bond, %V_{bur} of 6 varies from 31.8 (θ = 0°) to 28.5 (θ = 100°). Further rotation leads to an increase of the steric hindrance as the naphthyl ring is pointed toward the metal center. A maximum %V_{bur} of 43.8 is reached when at the dihedral angle of θ = 180°.

3.4. Topographic steric map of their metal complexes

Catalyst design has always been a challenging task and often driven by trial and error, or intuition, rather than a rational science. A classic solution to the problem is to use molecular descriptors capable of visualizing the catalysts space to offer a rational understanding of the designed catalyst. Using SambVca 2 tool, topographic steric map of the carbenes and, therefore, catalytic pocket of their metal complexes can be easily obtained.

For the six carbenes, when looking at the carbene from the metal center, along the M-C carbene bond (Figure 9a), topographic steric map of the carbenes appears as visualized in Figure 9b-g. The contours represent relative distance to the plane perpendicular to viewing axis and passing through the C_{carbene} atom.

Topographic steric map study reveals that the metal coordinated to NHC 1, 3, 6 are relatively accessible from the directions which is perpendicular to the carbene heterocycle planes. On the other hand, in the complex of 5, the catalytic pocket is relatively limited in size, and the incoming agent has to approach from direction opposite to carbene 5 to reach to the metal center. It is therefore suggested that the complex of 5 may not be the best choice of catalyst to activate bulky substrates.

4. Conclusion

Stereoelectronic properties of a series of 1,2,4-triazole-derived carbenes bearing different N4-substituents, namely isopropyl, benzyl, phenyl, mesityl, 2,6-diisopropylphenyl and 1-naphthyl, has been examined. The results suggest the similarity in nature and energy level of their HOMOs. Steric properties of the NHCs was evaluated and quantified using percent volume buried (%V_{bur}) methodology. The NHC with isopropyl N4-substituent was the least bulky one and the most bulky is the one with 2,6-diisopropylphenyl N4-substituent. Importantly, the NHCs with phenyl and 1-naphthyl N4-substituents display flexible steric properties, which were accessible by rotation of the phenyl or 1-naphthyl around the N-C single bond.

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