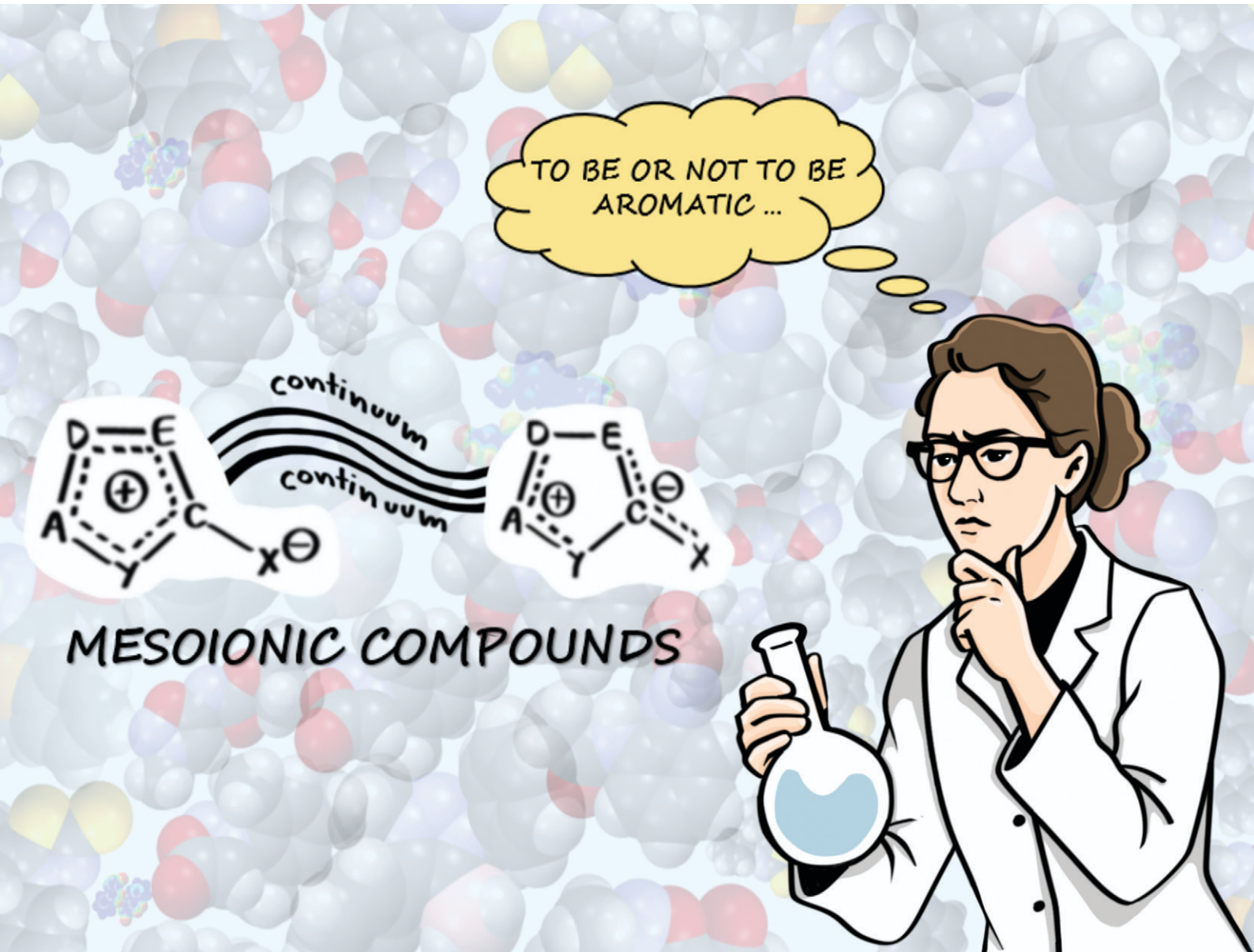


# CrystEngComm

[rsc.li/crystengcomm](http://rsc.li/crystengcomm)



ISSN 1466-8033



Cite this: *CrystEngComm*, 2025, 27, 4264

## Evaluation of mesoionic compound aromaticity using the HOMHED index†

Marcos Antonio Pinto Martins, <sup>\*a</sup> Tainára Orlando, <sup>b</sup>

Jéssica Maria Luis Rosa,

Priscila Santos Vieira de Lima <sup>a</sup> and Paulo Roberto dos Santos Salbego <sup>c</sup>

Mesoionic compounds are a class of heterocyclic compounds whose aromaticity has ignited considerable debate within the scientific community. This study aims to systematically evaluate the aromaticity of 25 mesoionic compounds across eight structural classes, utilizing experimental X-ray diffraction data in conjunction with the HOMHED geometric index to clarify this issue. The findings reveal a wide range of aromaticity (0.494–0.976), influenced by the type of heteroatoms in the heterocycle and exocyclic positions. Compounds containing oxygen atoms typically exhibit lower aromaticity, with HOMHED values below 0.7. In contrast, those containing nitrogen and boron atoms, especially in exocyclic positions, show higher aromaticity, with HOMHED values exceeding 0.9. Remarkably, compounds featuring exocyclic boron atoms displayed the highest aromaticity, with HOMHED values between 0.951 and 0.976. Compound 25 stood out with the lowest aromaticity, having a HOMHED value of 0.494, arising as a missing link, representing an important factor in elucidating the boundaries of the continuum of aromaticity. The study found that, on average, HOMHED values increase in the order of heterocyclic atom (Y): O1 < S1 < N1 and exocyclic atom (X): O6 < S6 ≈ N6 < B6. The research compares two canonical forms, the aromatic structure (AS) and the betaine structure (BS), to assess their impact on mesoionic compounds' overall resonance and aromatic character. Compounds with higher aromaticity demonstrated increased resonance in the D-A-Y and D-E-C-Y fragments, indicating a predominant AS contribution, whereas those with lower aromaticity showed a stronger BS influence. By introducing the AS index ( $I^{AS}$ ) and the BS index ( $I^{BS}$ ), this study quantitatively distinguishes the contributions of each structure. The findings challenge the binary view of mesoionic compounds as merely aromatic or non-aromatic, suggesting instead that these compounds exhibit varying degrees of aromaticity, as a continuum. The study concludes that the appropriate question is not whether mesoionic compounds are aromatic, but rather, 'How aromatic are mesoionic compounds?'.

Received 27th March 2025,  
Accepted 30th April 2025

DOI: 10.1039/d5ce00334b

rsc.li/crystengcomm

## Introduction

According to the International Union of Pure and Applied Chemistry (IUPAC), mesoionic compounds are dipolar heterocyclic compounds in which both the negative and positive charges are delocalized. These compounds, which cannot be adequately described by a totally covalent structure

nor represented satisfactorily by any single polar structure,<sup>1</sup> contain two or more heteroatoms. The formal positive charge is associated with the ring atoms, whereas the formal negative charge is linked to ring atoms or an exocyclic nitrogen or chalcogen atom.<sup>1</sup>

The structure of mesoionic compounds has attracted significant attention in structural chemistry and has spurred numerous theoretical studies aimed at elucidating their electronic structures.<sup>2–5</sup> Much of the theoretical work on mesoionic compounds focuses on discussing their aromaticity. While some researchers assert that mesoionic compounds are aromatic,<sup>6,7</sup> others firmly contend that they are not.<sup>4,8</sup>

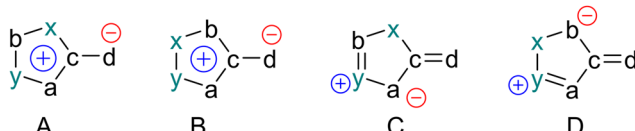
The debate over the aromaticity of mesoionic compounds stems from how these compounds are represented. The controversy began in 1955 when Kutzinsky urged the scientific community to refrain from using the term 'mesoionic

<sup>a</sup> Núcleo de Química de Heterociclos (NUQUIMHE), Department of Chemistry, Federal University of Santa Maria (UFSM), 97105-900, Santa Maria, RS, Brazil.  
E-mail: martins.marcos@uol.com.br

<sup>b</sup> Academic Department of Chemistry, Federal Technological University of Paraná (UTFPR), 85884-000 Medianeira Campus, PR, Brazil

<sup>c</sup> Núcleo de Química de Heterociclos (NUQUIMHE), Departamento de Engenharia e Tecnologia Ambiental (DETA), Federal University of Santa Maria (UFSM), 98400-000, Frederico Westphalen, RS, Brazil

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d5ce00334b>



Scheme 1 Structures used to represent mesoionic compounds.

compounds' and to instead refer to them as 'mesomeric betaines'.<sup>9</sup> This nomenclature persisted until 1985 when Ollis, Stanforth, and Ramsden introduced a new classification based on the different types of heterocycle conjugations (Scheme 1).<sup>2</sup>

Subsequently, mesoionic compounds were categorized into two types based on their heteroatoms' arrangement, with A and B serving as positional isomers. The tautomerism in structures A and B, showcasing the heterocycle as aromatic, has been utilized to explore, for example, the reactivity of some mesoionic compounds (Scheme 1).<sup>2,3</sup>

Although the representation of forms A and B has been widely recommended for years, it has been consistently questioned due to its suggestion of aromatization in the heterocycle. Despite the conjugated system adhering to Hückel's rule, the aromatic nature of these heterocyclic rings has always been questioned. For instance, the reactivity of mesoionic compounds towards electrophiles (SnAr) could suggest their aromatic character;<sup>10</sup> however, their ability to react as 1,3-dipoles presents evidence against such aromatic behavior.<sup>11</sup>

In pioneering work, Simas *et al.*<sup>8</sup> in 1998 concluded that mesoionic compounds are not aromatic, based on X-ray diffraction data and theoretical studies, which indicate clearly separated positive and negative regions functioning independently as electron-deficient and electron-rich areas. Following this, Champagne and Houk<sup>4</sup> proposed structures C and D for mesoionic compounds, relying on theoretical data and referencing X-ray data from Simas,<sup>8,12–14</sup> although they overlooked critical experimental observations about the presence of two electron-delocalized regions, calling into question the validity of structures C and D.

Conversely, M. Duvall in 2013<sup>6</sup> demonstrated through extensive theoretical analysis using Bird and nucleus independent chemical shift (NICS) aromaticity indices that mesoionic compounds, including sydnone and münnchnone, are aromatic, exhibiting indices comparable to furans and pyrroles. Similarly, Kuroda *et al.*<sup>7</sup> based on experimental X-ray diffraction data contributed to the general idea of aromaticity of mesoionic compounds, demonstrating that these compounds possess a moderate degree of aromaticity, showing that the X-ray diffraction data aligned consistently with previous computational predictions.

More recently, considering the research conducted by several scientists, including Champagne and Houk,<sup>4</sup> it was determined that the bond, charge, and magnetic field data in the heterocyclic ring indicated a lack of aromatization in mesoionic compounds. Therefore, structures C and D (Fig. 1) were recommended for representing mesoionic compounds.

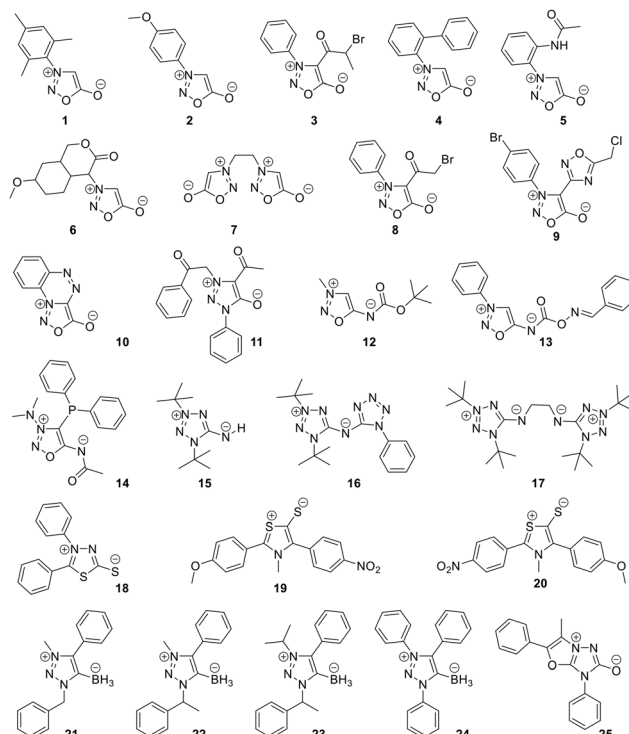


Fig. 1 Mesoionic compounds 1–25 used in this study.

Following Champagne and Houk's<sup>4</sup> research, Porte *et al.* published a review in 2021,<sup>5</sup> adopting and recommending the presentation of mesoionic compounds as structures C and D. Their recommendation was based on a theoretical study of cycloaddition reactions<sup>4</sup> and did not include data on aromaticity indices, merely referencing the work by Simas *et al.*<sup>8</sup> and Katritzky *et al.*<sup>15</sup> as though these articles encompassed all known aromaticity data for mesoionic compounds.

From these controversial positions, it is clear that the aromaticity of mesoionic compounds remains a contested issue. In response, we embarked on this study to systematically investigate the aromaticity of mesoionic compounds and contribute to the ongoing debate: are mesoionic compounds aromatic? To this end, we employed experimental X-ray data alongside the HOMHED geometric index<sup>16</sup> to evaluate the aromaticity of these compounds. Our study covers a collection of 25 mesoionic compounds across eight distinctive classes, offering diverse structures for analysis (Fig. 1).

## Experimental

### X-ray structures

The X-ray data for mesoionic compounds numbered 1–25 used in this study were collected from the CCDC database using the software Conquest (CSD version 5.45). The bond lengths (in Ångströms) are listed in Table S1.† The search involved drawing the specific nuclei of mesoionic structures. Subsequently, a manual search was conducted to identify the

structures with the best crystallographic data, resulting in 25 structures. Crystallographic information files (CIF) for compounds 1–25 are deposited at the Cambridge Crystallographic Data Centre (CCDC) under the following identification refcodes: YOMWAA (1),<sup>17</sup> WADNOG (2),<sup>18</sup> JOMBUK (3),<sup>19</sup> EWOXAP (4),<sup>20</sup> SOLZOJ (5),<sup>21</sup> BIJKEN (6),<sup>22</sup> ETHBSD (7),<sup>23</sup> XAZTOJ (8),<sup>24</sup> XAZJEQ (9),<sup>25</sup> XZBTZO (10),<sup>26</sup> IJOWUB (11),<sup>27</sup> DEWYIQ (12),<sup>28</sup> TUFPIW (13),<sup>29</sup> KUCYEN (14),<sup>30</sup> OVIWAU (15),<sup>31</sup> OVIWIC (16),<sup>31</sup> OVIWEY (17),<sup>31</sup> RIPROB (18),<sup>32</sup> MAGTEV (19),<sup>33</sup> MAGTIZ (20),<sup>33</sup> KIXHUW (21),<sup>34</sup> KIXJAE (22),<sup>34</sup> KIXJEI (23),<sup>34</sup> KIXJIM (24),<sup>34</sup> and ZOCLAI (25).<sup>7</sup>

### HOMHED index

The HOMHED index was used to calculate the aromaticity of mesoionic compounds' structures.<sup>16</sup> To calculate the HOMHED index, it was necessary to obtain the  $R_{\text{opt}}$  bond length and the normalized constant ( $\alpha$ ) for each type of bond. The  $R_{\text{opt}}$  bond length was calculated according to eqn (1), where  $R_s$  and  $R_d$  are the reference single and double bond lengths, respectively, taken from experimental X-ray data, respectively, and  $\omega$  is the ratio of stretching force constants for pure double and single bonds. The  $\omega$  ratio close to 2 was used for the C–C and C–X bonds. The normalized constant ( $\alpha$ ) for each type of bond was calculated using eqn (2). The HOMHED, a dimensionless parameter, is calculated for each compound using eqn (3), where  $n$  is the number of bonds considered in the resonance system, and  $R_i$  represents the bond lengths. Table 1 presents the data for  $R_s$ ,  $R_d$ , and  $R_{\text{opt}}$  for C–C, C–N, C–O, C–S, N–N, N–O, and N–S bonds based on data from the literature.<sup>16</sup>

$$R_{\text{opt}} = \frac{(R_s + \omega R_d)}{(1 + \omega)} \quad (1)$$

$$\alpha = \frac{2}{(R_{\text{opt}} - R_s)^2 + (R_{\text{opt}} - R_d)^2} \quad (2)$$

$$\text{HOMHED} = 1 - \frac{\alpha}{n} \sum (R_{\text{opt}} - R_i)^2 \quad (3)$$

### Bond lengths for HOMHED

The bond lengths for HOMHED are detailed in Table 1. Data for  $R_s$ ,  $R_d$ , and  $R_{\text{opt}}$ , related to C–C, C–N, C–O, C–S, N–N, N–O,

and N–S bonds, were derived from Frizzo and Martins.<sup>16</sup> The C–N<sup>+</sup> and C–B bond lengths were determined using eqn (1) and (2) based on the data below.

**C–N<sup>+</sup> bond lengths.** The bond lengths for pure single ( $R_s$ ) and double ( $R_d$ ) bonds in non-conjugated systems, or systems without  $\pi$ -electron and/or n-electron delocalization, were obtained from X-ray data of compounds:  $\text{Me}_4\text{N}^+\text{I}^-$  (QQQCVG01),  $\text{Me}_4\text{N}^+\text{I}^-$  (QQQCVG02),  $\text{Me}_4\text{N}^+\text{Cl}^-$  (ZZZUQM02),  $\text{Me}_4\text{N}^+\text{Br}^-$  (ZZZUQO03),  $\text{Me}_4\text{N}^+\text{Br}^-$  (ZZZUQO04),  $\text{Me}_2\text{N}^+=\text{CH}_2\text{Br}^-$  (LILLOH),  $\text{Me}_2\text{N}^+=\text{CHO}_2\text{Cl}^-$  (VAPREJ), and  $\text{Et}_2\text{N}^+=\text{CH}_2\text{Cl}^-$  (TIHGOH).<sup>35–42</sup>

**C–B bond lengths.** The bond lengths for pure single ( $R_s$ ) and double ( $R_d$ ) bonds in non-conjugated systems, or systems without  $\pi$ -electrons and/or n-electron delocalization, were identified using X-ray data for compounds with refcodes C–B (JAZGEX, SEHYEN)<sup>43,44</sup> and C= B (VARJED).<sup>45</sup>

## Results and discussion

Mesoionic compounds cannot be represented by a single contributing structure among several canonical forms described in various mesomeric structures and sometimes suggest a high level of aromaticity in their heterocycles. We propose that a mesoionic compound, regardless of the ring heteroatoms or the exocyclic atom, should be represented by two main canonical structures: the aromatic (AS) and betaine structures (BS), as shown in Scheme 2. In the AS, the formal positive charge of the mesoionic compound is associated with all the ring atoms (D–A–Y–C–E), and a formal negative charge is associated with the exocyclic atom (X). In the BS, the formal positive charge of mesoionic compounds is associated with the ring atoms (D–A–Y), and the formal negative charge is associated with both ring atoms and an exocyclic atom (E–C–X). Going forward, we aim to bridge the gap in research on mesoionic compounds' aromaticity by seeking evidence to determine which canonical structures (*i.e.*, AS or BS) best represent the mesoionic compounds' structures.

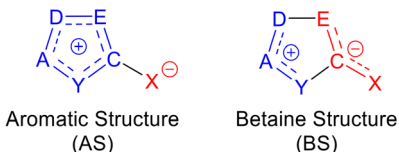
### The HOMHED index

Initially, we applied the HOMHED index to evaluate the aromaticity of 25 mesoionic compounds. Introduced in 2012 by Frizzo and Martins,<sup>16</sup> it builds on the original HOMA concept for assessing the aromaticity of heterocycles<sup>46</sup> and utilizes experimental single-crystal X-ray diffraction data to estimate optimal bond lengths, classifying compounds based

**Table 1** Single ( $R_s$ ), double ( $R_d$ ), optimal ( $R_{\text{opt}}$ ) bond lengths, and the normalized constants ( $\alpha$ ) used in the HOMHED calculations

Bond	C–C	C–N	C–O	C–S	N–N	N–O	N–S	C–B	C–N <sup>+</sup>
$R_s$ (Å)	1.530	1.474	1.426	1.819	1.454	1.463	1.765	1.600	1.494
$R_d$ (Å)	1.316	1.271	1.210	1.599	1.240	1.218	1.541	1.362	1.266
$R_{\text{opt}}$ (Å)	1.387	1.339	1.282	1.672	1.311	1.300	1.616	1.441	1.342
$\alpha$	78.6	87.4	77.2	74.4	78.6	60.0	71.7	63.4	69.3
Reference	16	16	16	16	16	16	16	[This study] <sup>a</sup>	[This study] <sup>a</sup>

<sup>a</sup> Data determined using eqn (1) and (2).



**Scheme 2** The canonical aromatic and betaine structures of mesoionic compounds.

on how closely their values approach 1, with those closer to 1 being deemed more aromatic. This index has been used for a variety of five- and six-membered heterocycles with C–C, C–N, C–O, C–S, N–N, N–O, and N–S bonds.<sup>16</sup> A comprehensive review on the geometric criteria for aromaticity and the development of HOMA-based indices was published by Krygowski *et al.*<sup>47</sup>

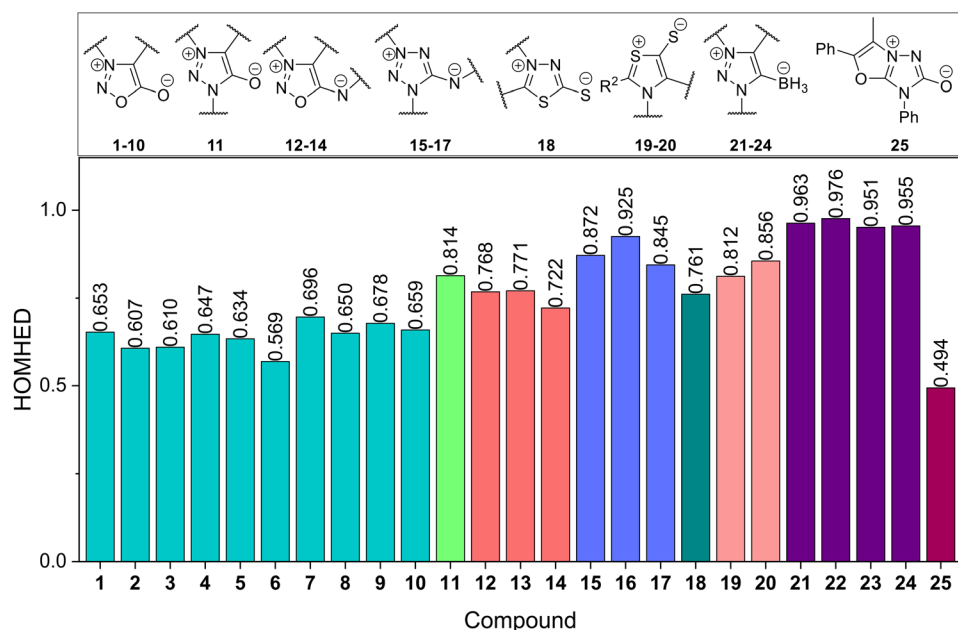
Nevertheless, to encompass the full range of compounds in this study, it became necessary to determine the optimal bond length ( $R_{\text{opt}}$ ) for C–B bonds; the  $R_{\text{opt}}(\text{C–B})$  values were determined using the same methodology<sup>16</sup> (Table 1). The optimal bond lengths  $R_s$ ,  $R_d$  were derived from experimental X-ray data of compounds containing both C–B and C=≡B bonds. Despite Zborowski *et al.*<sup>48</sup> previous publication of  $R_{\text{opt}}$  for the C–B bond based on X-ray diffraction data, we observed that the authors used a C<sup>sp<sup>3</sup></sup>–B<sup>sp</sup> bond for  $R_s$ . This single bond is shorter than that of a ‘pure’ C<sup>sp<sup>3</sup></sup>–B<sup>sp<sup>2</sup></sup> bond by approximately 0.05 Å, leading to an underestimated  $R_{\text{opt}}$  (1.424 Å) and an overestimated  $\alpha$  (104.5). Conversely, aiming to use the same criteria in our 2012 work,<sup>16</sup> we used X-ray diffraction data from ‘pure’ single bonds of compounds containing C<sup>sp<sup>3</sup></sup>–B<sup>sp<sup>2</sup></sup> bond to determine the  $R_{\text{opt}}(\text{C–B})$ .

The HOMHED indices for compounds 1–25 were calculated using eqn (3), which considers the bond lengths

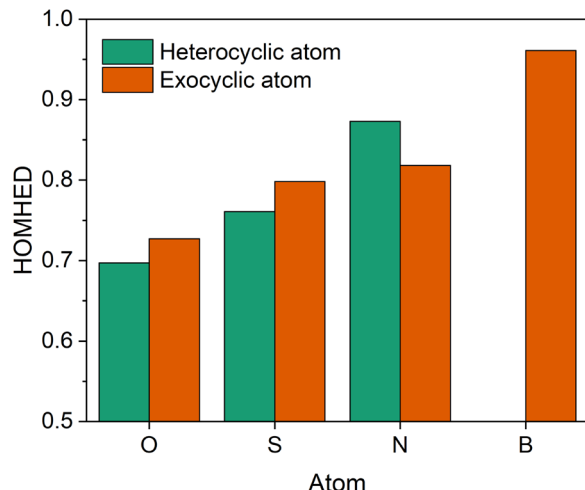
( $R_i$ ) between atoms Y–A, A–D, D–E, E–C, C–Y in the heterocyclic ring and C–X (exocyclic bond) (Table S1 in ESI†). Compound 25 has the lowest aromaticity index (0.494), while compounds 21–24 show the highest aromaticity indices (0.951–0.976). Fig. 2 illustrates the distribution of HOMHED data for compounds 1–25.

By observing the HOMHED values of the compounds, we see that compounds 1–10 exhibit low aromaticity (0.569–0.696), as expected for heterocycles containing an oxygen atom (O1) in the ring. This phenomenon, where a cyclic oxygen atom promotes lower aromaticity, has already been reported.<sup>49,50</sup> Compound 11 displays an increase in aromaticity due to the substitution of O1 with N1 in the heterocyclic ring (HOMHED = 0.8). In compounds 12–14, the HOMHED values vary at 0.72–0.77; this is related to the replacement of the exocyclic oxygen atom with a nitrogen atom; however, unlike the previous example, the increase in aromaticity was very subtle. In both scenarios, substituting a chalcogen atom for a nitrogen atom results in an increase in HOMHED, suggesting that mesoionic compounds containing oxygen atoms, whether in the heterocycle or exocyclicly, have lesser aromaticity.

Mesoionic tetrazolium derivatives (compounds 15–17) and thiazole derivatives (compounds 19–20) also exhibit HOMHED values above 0.8. Compound 18 has a lower HOMHED than compounds 15–17 and 19–20, likely due to the replacement of N1 with S1. Lastly, compounds 21–24 present HOMHED values around 0.95, suggesting an aromaticity similar to that of uncharged nitrogenous heterocyclic compounds.<sup>16</sup> This can be attributed to the fact that the negative charge on the exocyclic boron atom does not resonate with the heterocyclic ring, unlike in compounds 1–20. In summary, the average HOMHED values increase in



**Fig. 2** HOMHED values of compounds 1–25. The colored bars represent the different structures of mesoionic compounds.



**Fig. 3** Effect of heterocyclic (Y) and exocyclic (X) atom on the aromaticity of mesoionic compounds 1–24.

the order of heterocyclic atom (Y): O1 < S1 < N1; and for the exocyclic atom (X): O6 < S6 ≈ N6 < B6, as shown in Fig. 3.

Although Kuroda *et al.*<sup>7</sup> described compound 25 and calculated its HOMHED and NICS indices, they concluded that the synthesized mesoionic triazolone exhibits a moderate degree of aromatic character. The authors did not present a comparative study to demonstrate how this type of structure's HOMHED index compares to those of other mesoionic compounds with similar structures. Our comparative study indicates that compound 25 may be an exception, as it has a very low HOMHED value (0.494). Therefore, arising as a missing link, representing an important factor in elucidating the boundaries of the aromaticity continuum. For 5-membered heterocycles containing a  $sp^3$ -nitrogen in the ring, HOMHED values greater than 0.800 are typically expected. This suggests that other factors may decrease the aromaticity of the heterocycle, with the most significant effect likely being the tension between the two condensed 5-membered rings that comprise the molecule.

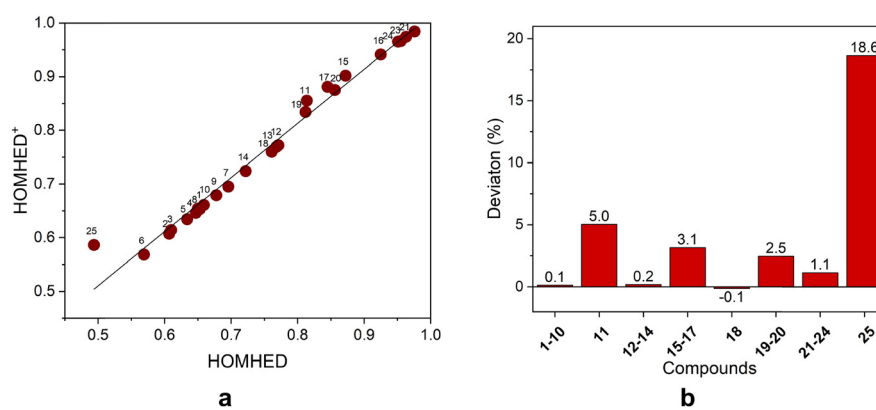
### The HOMHED vs. HOMHED<sup>+</sup>

The next issue to address concerns determining the HOMHED of aromatic compounds containing charged nitrogen, such as mesoionic or salts. More specifically, would it be more accurate to calculate the HOMHED value with the  $R_{opt}(C-N)$  bond or with the  $R_{opt}(C-N^+)$  bond? To answer this question, this study presents, for the first time in the literature, the optimal bond ( $R_{opt}$ ) values for C–N<sup>+</sup>. The HOMHED<sup>+</sup> data for compounds 1–25, which utilize  $R_{opt}(C-N^+)$  instead of  $R_{opt}(C-N)$ , can be found in Table S2 (ESI†). Overall, the results show no significant differences between the HOMHED and HOMHED<sup>+</sup> values, whether using the  $R_{opt}(C-N)$  or the  $R_{opt}(C-N^+)$  bond (Fig. 4a).

In the series of compounds studied, the use of  $R_{opt}(C-N^+)$  increases HOMHED<sup>+</sup> value by a range of 0–5%, depending on the number of C–N bonds in the heterocycle and the presence of nitrogen-sp<sup>3</sup> ( $\pi$ -excess heterocycles), as shown in Fig. 4b. A notable exception is the mesoionic compound 25, which consists of two fused 5-membered rings with 4 C–N bonds. For this compound, the HOMHED<sup>+</sup> value is 18% higher than HOMHED. Furthermore, the application of  $R_{opt}(C-N^+)$  bonds in certain tetrazolium<sup>+</sup>, imidazolinium<sup>+</sup>, and pyridinium<sup>+</sup> salts was evaluated. The results showed that the HOMHED<sup>+</sup> value was over 2% higher for imidazolinium<sup>+</sup> salts, while it was negligible for the other two series of salts.

### Aromatic vs. betaine structures

Generally, researchers have posited that mesoionic compounds consist of a five-member heterocyclic ring in which the D–A–Y atoms carry the positive charge, while the E–C–X atoms carry the negative charge, similar to betaines. In contrast, another form presents the positive charge distributed across the D–A–Y–C–E atoms of the ring, with the negative charge localized on the exocyclic X atom, resembling aromatic compounds, as illustrated in Fig. 5. This distinction underpins the understanding that the lower aromaticity observed in mesoionic compounds is attributed to the betaine-like structure. This structure significantly reduces the



**Fig. 4** (a) Correlation between HOMHED and HOMHED<sup>+</sup>;  $R = 0.988$  and (b) variation of the HOMHED index for compounds 1–25. Calculated by  $((HOMHED^+/HOMHED) - 1) \times 100$ .

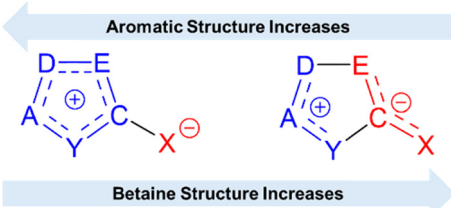


Fig. 5 Charge separation in the canonical aromatic and betaine structures of mesoionic compounds.

resonance in the Y-C and D-E bonds, limiting the interaction between the positive and negative charge resonance systems.<sup>8</sup>

In this context, considering the betaine and aromatic forms, it seems important to investigate the resonance (electronic density delocalization) in fragments that comprise the betaine structure: D-A-Y, E-C-X fragments, and the D-E, C-Y fragments that connect D-A-Y and E-C-X. Our preliminary understanding suggests that a mesoionic compound exhibits several canonical structures, notably the AS and BS, each contributing to varying extents to the mesoionic compound's properties (Fig. 5). Thus, we cannot prefer one canonical structure over another but rather need to understand the extent of each structure's contribution.

Using HOMHED, it is possible to calculate the resonance for selected fragments, that is, not to the entire molecule, but to specific fragments of the analyzed structure.<sup>16</sup> This method allowed us to obtain the resonance data for the residues  $\text{Res}_{(\text{ECX})}$ ,  $\text{Res}_{(\text{DAY})}$ ,  $\text{Res}_{(\text{DE,CY})}$  in compounds 1–25 (Table 2). With this data, we aim to determine how the aromatic character of a mesoionic compound correlates with the resonance of its fragments.

With resonance data available, analyzing the mesoionic compound's aromatic character—in terms of greater or lesser—based on the HOMHED geometric data,<sup>16</sup> relies on correlating it with the resonances of the fragments (D-A-Y), (D-E, C-Y), and (E-C-X). In this context, we anticipate: (i) high resonance indices for the fragments (D-A-Y) and (D-E, C-Y)

and low resonance indices for the fragment (E-C-X) in compounds with high aromaticity indices (HOMHED); and (ii) compounds exhibiting high resonance indices in the fragments (D-A-Y) and (E-C-X) will display characteristics typical of betaines. The resonance index in the fragment (D-E, C-Y) will determine the extent to which the resonances of the two fragments (D-A-Y) and (E-C-X) differ, predominantly contributing to the betaine characteristic. We associate the HOMHED index with fragment resonances to gather additional insights into mesoionic behavior (Fig. 6). The data points on the graph are categorized into three primary regions.

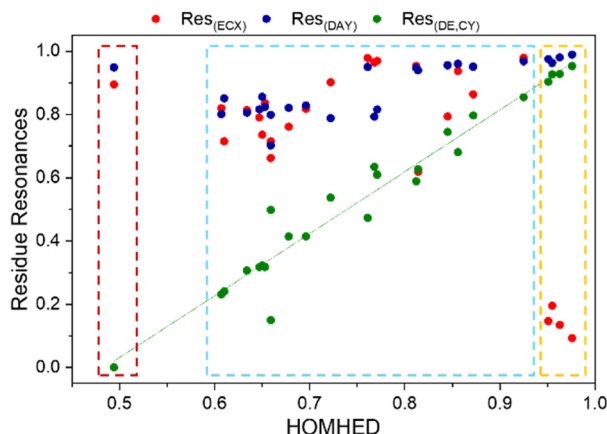
As shown in Fig. 6, in region A, the resonance data  $\text{Res}_{(\text{DAY})}$ ,  $\text{Res}_{(\text{DE,CY})}$ , and  $\text{Res}_{(\text{ECX})}$  for compounds 21–24 are presented, showcasing HOMHED values greater than 0.950. It is evident that  $\text{Res}_{(\text{DAY})}$  and  $\text{Res}_{(\text{DE,CY})}$  values exceed 0.9, while  $\text{Res}_{(\text{ECX})}$  values fall below 0.2. This indicates that compounds 21–24 possess a degree of aromaticity and highlight the minimal contribution of the canonical betaine structure. It is important to note that these compounds contain a boron atom carrying a negative charge, which does not resonate with the ring atoms, as reflected by the low  $\text{Res}_{(\text{ECX})}$  values.

Region B displays the highest values of  $\text{Res}_{(\text{DAY})}$  and  $\text{Res}_{(\text{ECX})}$ , although the lowest of  $\text{Res}_{(\text{DE,CY})}$ . Compound 25, the sole compound in this region, exhibits values of  $\text{Res}_{(\text{DAY})} = 0.985$ ,  $\text{Res}_{(\text{ECX})} = 0.949$ , and  $\text{Res}_{(\text{DE,CY})} = 0$ , with a HOMHED of 0.494. These figures suggest compound 25 has a low aromaticity level and a significant contribution from the canonical betaine structure.

In region C, the resonance data  $\text{Res}_{(\text{DAY})}$  and  $\text{Res}_{(\text{ECX})}$  for most studied compounds presenting fragment resonance greater than 0.7 and HOMHED values ranging from 0.60 to 0.95 are found. This region may indicate a notable contribution of the canonical betaine structure, although the  $\text{Res}_{(\text{DE,CY})}$  data span a wide range (0.150–0.855), pointing out the need for additional data to more accurately quantify the contributions of each canonical aromatic or betaine structure.

Table 2 Resonances of the fragments  $\text{Res}_{(\text{DAY})}$ ,  $\text{Res}_{(\text{DE,CY})}$  and  $\text{Res}_{(\text{ECX})}$  of compounds 1–25

Compound	$\text{Res}_{(\text{ECX})}$	$\text{Res}_{(\text{DAY})}$	$\text{Res}_{(\text{DE,CY})}$	Compound	$\text{Res}_{(\text{ECX})}$	$\text{Res}_{(\text{DAY})}$	$\text{Res}_{(\text{DE,CY})}$
1	0.836	0.824	0.317	14	0.902	0.788	0.537
2	0.820	0.801	0.232	15	0.864	0.951	0.797
3	0.715	0.851	0.241	16	0.979	0.968	0.855
4	0.790	0.816	0.317	17	0.794	0.956	0.745
5	0.814	0.806	0.306	18	0.978	0.950	0.473
6	0.715	0.799	0.150	19	0.953	0.947	0.589
7	0.818	0.829	0.414	20	0.938	0.960	0.680
8	0.736	0.856	0.322	21	0.135	0.980	0.929
9	0.761	0.821	0.414	22	0.093	0.989	0.953
10	0.662	0.702	0.498	23	0.147	0.975	0.904
11	0.619	0.940	0.627	24	0.196	0.963	0.927
12	0.965	0.793	0.634	25	0.895	0.949	0.000
13	0.969	0.816	0.610				

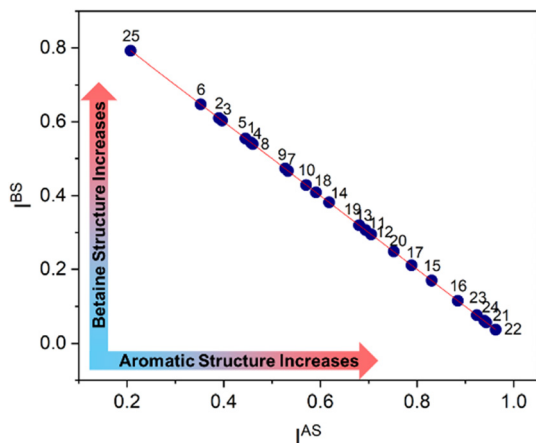


**Fig. 6** Correlation of  $\text{Res}_{(\text{DAY})}$ ,  $\text{Res}_{(\text{DE,CY})}$ , and  $\text{Res}_{(\text{ECX})}$  parameters with HOMHED. A region is in the orange box, B region is in the red box, and C region is in the blue box.

A critical observation is that the data  $\text{Res}_{(\text{DE,CY})}$  exhibits a strong linear correlation with the HOMHED data ( $r = 0.973$ ;  $N = 25$ ; equation:  $\text{Res}_{(\text{DE,CY})} = 1.958 \text{ HOMHED} - 0.947$ ), indicating the varying resonance in the D-E and C-Y bonds. High  $\text{Res}_{(\text{DE,CY})}$  values suggest a stronger aromatic character in the mesoionic compounds. This fact shows that the degree of aromaticity can be determined quantitatively from an error calculation between the HOMHED and  $\text{Res}_{(\text{DE,CY})}$  in relation to 1, as demonstrated by eqn 4, which provides the aromatic structure index ( $I^{\text{AS}}$ ). Eqn (5) provides the betaine structure index ( $I^{\text{BS}}$ ) of mesoionic compounds. The  $I^{\text{AS}}$  and  $I^{\text{BS}}$  data for the 25 compounds are listed in Table S3 (ESI†) and shown in Fig. 7.

$$I^{\text{AS}} = 1 - \sqrt{\frac{(1 - \text{HOMHED})^2 + (1 - \text{Res}_{(\text{DE,CY})})^2}{2}} \quad (4)$$

$$I^{\text{BS}} = 1 - I^{\text{AS}} \quad (5)$$



**Fig. 7** Aromatic structure index ( $I^{\text{AS}}$ ) and betaine structure index ( $I^{\text{BS}}$ ) for compounds 1–24.

Considering the average HOMHED of each class of mesoionic compounds (1–10, 11, 12–14, 15–17, 18, 19–20, 21–24, 25), we compared them with the average HOMHED of a series of different heterocycles that have been previously published (Fig. 8).<sup>16</sup> This comparison allows us to observe that the HOMHED increases from the least aromatic compounds, which are the 5-membered rings containing oxygen (e.g., oxazoles, furans, isoxazoles), to the sulfur-containing compounds (e.g., thiophenes, thiazoles), then to the nitrogenous compounds (e.g., imidazoles, pyrazoles, pyrroles), and finally to the nitrogenous compounds with 6-membered rings (e.g., pyrimidines, pyridines, pyrazines).

In Fig. 8, one can observe that mesoionic compounds 1–24 are distributed based on HOMHED values in the same order as uncharged heterocycles with similar structures. The three series of charged heterocycles, namely, the imidazolium, tetrazolium, and pyrimidinium salts, follow the same order as the uncharged heterocycles. However, an exception was noted with compound 25, which exhibits a HOMHED value lower than expected compared to 5-membered rings containing an  $\text{N-}^{\text{sp}^3}$ , typically characterized by HOMHED values greater than 0.800.

Previous research on this topic of mesoionic aromatic order includes a study by Bird<sup>50</sup> demonstrating that the introduction of an oxygen atom into the ring decreases the aromaticity of heterocycles. Frizzo and peers,<sup>16</sup> in their studies on the aromaticity of heterocyclic compounds, found an increase in aromaticity in the sequence furan → thiophene → pyrrole. Morzherin *et al.*<sup>51</sup> showed that mesoionic compounds, such as 1,2,3-triazoles and 1,2,3,4-tetrazoles, are more aromatic than other mesoionic compounds, considering them more as ‘masked dipoles’. Moreover, Frizzo *et al.*<sup>16</sup> also showed that 5- and 6-membered heterocyclic compounds that contain a larger number of nitrogen atoms are more aromatic than their less nitrogenous counterparts.

Our preliminary study suggests that the aromaticity of heterocycles is minimally impacted by the presence of charges within the heterocycle, whether in mesoionic compounds (1–24) or salts (imidazolinium $^+$ , tetrazolinium $^+$ , pyridinium $^+$ ), especially when evaluating the aromaticity order among 5- or 6-membered heterocycles. This phenomenon of ‘charges not mattering’ has recently been reported in a recent study by our group<sup>32</sup> covering supramolecular behavior of mesoionic compounds 1–24 based on their X-ray diffraction data. The findings revealed that the mesoionic compounds exhibit behavior similar to that of uncharged compounds.<sup>32</sup>

Furthermore, the supramolecular data indicated that these compounds exhibit characteristics typical of aromatic compounds because (i) the molecular electrostatic potential predominantly shows positive potential in the ring and negative potential in the exocyclic heteroatom; (ii) there are numerous intermolecular interactions based on the stacking of mesoionic rings with the exocyclic atom; (iii) stackings between mesoionic rings and phenyls are also observed; and

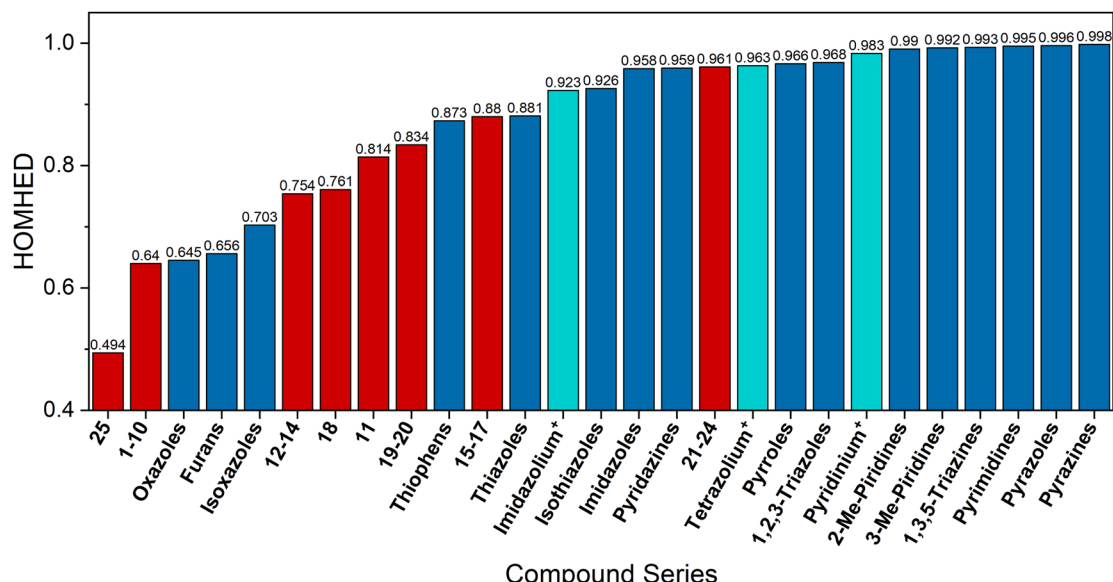


Fig. 8 Comparison of the average HOMHED of the series of mesoionic compounds 1–25 (red), heterocyclic salts (cyan), and heterocycles (blue).

(iv) most of the crystallization mechanisms of the studied mesoionic compounds involve molecular stacking structures of the 1D nuclei and 1D-dimer types.<sup>32</sup>

## Conclusions

This study thoroughly investigated the aromaticity of 25 mesoionic compounds, divided into eight structural categories, by utilizing the HOMHED geometric index and experimental X-ray diffraction data. The combination of experimental and theoretical approaches contributes significantly to the debate on the aromatic nature of mesoionic compounds and refines the understanding of their electronic structures. This understanding has implications for their reactivity, stability, and potential applications in supramolecular chemistry. The findings reveal a substantial variation in aromaticity among the compounds, primarily influenced by the nature of the heteroatoms and exocyclic atoms. This underscores the necessity for a nuanced approach to evaluating the electronic properties of mesoionic compounds beyond the oversimplified definition of to be or not to be aromatic.

Our findings revealed that the degree of aromaticity in mesoionic compounds varies depending on the heteroatoms present. Compounds containing oxygen atoms in the ring generally exhibit lower aromaticity, whereas those with nitrogen atoms, particularly when nitrogen is part of an exocyclic element or  $sp^3$  bond, tend to have higher aromaticity. Compounds 21–24 showed the highest aromaticity due to the presence of a negatively charged boron atom that does not resonate with the ring atoms.

Furthermore, this study highlights the significance of the aromatic and betaine structures in their canonical forms in understanding the aromatic nature of mesoionic compounds. Analysis of resonance data across different fragments within

the compounds revealed that high resonance in D-E and C-Y bonds correlates with increased aromatic character, whereas high resonance in D-A-Y and E-C-X fragments suggests a predominant betaine character. The introduction of the aromatic structure index and betaine structure index provides a quantitative method to assess the contributions of these canonical structures. Compounds such as 21–24 were found to predominantly exhibit aromatic character with minimal betaine contribution, while compound 25 displayed a dominant betaine structure, leading to its lower aromaticity. Additionally, our findings showed that mesoionic compounds 1–24 possess aromaticity comparable to uncharged heterocyclic compounds, indicating that their aromaticity is minimally impacted by the presence of charges within the heterocycle.

It is often mistakenly accepted that mesoionic compounds are not aromatic, a belief that has become widespread. This suggests the need for a critical perspective devoid of preconceived notions, as the binary question of “to be or not to be aromatic” may be oversimplifying the matter. Focusing on one canonical structure over another due to formal charges in mesoionic compounds can lead to a skewed view. While research on mesoionic compounds has contributed valuable insights into synthesis, reactivity, and structure, it often overlooks a detailed analysis of aromaticity. Many investigations into their non-aromatic nature are based solely on theoretical data, creating a potential risk. These results can contain circular data output; the data introduced into the systems and the response data are part of the same type of calculation, with no experimental data breaking this theoretical bubble created.

Nevertheless, from a systematic perspective, with the researcher suspending any preconceptions, it is possible to observe that mesoionic compounds are heterocycles with all Hückel's requirements to behave with some degree of

aromaticity. Regardless of the index used for measuring, most mesoionic compounds in the literature display a wide range of aromaticity indices comparable to those of heterocycles such as furans and thiophenes. Hence, mesoionic compounds are aromatic, though the extent of their aromaticity still needs clarification. Therefore, supported by experimental data, we propose that the correct scientific question to be answered here is: how aromatic are mesoionic compounds?

## Data availability

The data that support the findings of this study are available in the ESI† of this article.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors acknowledge the Rio Grande do Sul Research Support Foundation (FAPERGS process no. 17/2551-0000981-9), the National Council for Scientific and Technological Development (CNPq process no. 405071/2016-7), and the Coordination for the Improvement of Higher Education Personnel (CAPES) for financial support. The fellowships from CNPq (MAPM) and CAPES (JMLR and PSVL) are also acknowledged.

## References

- 1 Mesoionic compounds, in *IUPAC Compendium of Chemical Terminology*, International Union of Pure and Applied Chemistry, 5th edn, 2025, DOI: [10.1351/goldbook.M03842](https://doi.org/10.1351/goldbook.M03842).
- 2 W. D. Ollis, S. P. Stanforth and C. A. Ramsden, *Tetrahedron*, 1985, **41**, 2239–2329.
- 3 C. A. Ramsden, *Tetrahedron*, 2013, **69**, 4146–4159.
- 4 P. A. Champagne and K. N. Houk, *J. Org. Chem.*, 2017, **82**, 10980–10988.
- 5 K. Porte, M. Riomet, C. Figliola, D. Audisio and F. Taran, *Chem. Rev.*, 2021, **121**, 6718–6743.
- 6 M. B. Duvall, *Master of Science thesis*, University of Georgia, 2013.
- 7 Y. Kuroda, M. Krell, K. Kurokawa and K. Takasu, *Chem. Commun.*, 2024, **60**, 1719–1722.
- 8 A. M. Simas, J. Miller and P. F. de Athayade Filho, *Can. J. Chem.*, 1998, **76**, 869–872.
- 9 A. J. Boulton, *Biogr. Mem. Fellows R. Soc.*, 2015, **61**, 225–245.
- 10 B. V. Badami, *Resonance*, 2006, **11**, 40–48.
- 11 W. P. Oziminski and C. A. Ramsden, *Tetrahedron*, 2015, **71**, 7191–7198.
- 12 K. K. Cheung, S. Galembeck, J. Miller, M. B. de Oliveira, A. B. Pereira and A. M. Simas, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1991, **47**, 2630–2632.
- 13 K. K. Cheung, S. Galembeck, J. Miller, M. B. de Oliveira, A. B. Pereira and A. M. Simas, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1992, **48**, 523–525.
- 14 K. K. Cheung, A. Echevarria, S. Galembeck, M. A. M. Maciel, J. Miller, V. M. Rumjanek and A. M. Simas, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1992, **48**, 1471–1474.
- 15 A. R. Katritzky, K. Jug and D. C. Oniciu, *Chem. Rev.*, 2001, **101**, 1421–1450.
- 16 C. P. Frizzo and M. A. P. Martins, *Struct. Chem.*, 2012, **23**, 375–380.
- 17 S. Wiechmann, T. Freese, M. H. H. Drafz, E. G. Hübner, J. C. Namyslo, M. Nieger and A. Schmidt, *Chem. Commun.*, 2014, **50**, 11822–11824.
- 18 H. K. Fun, J. H. Goh, Nithinchandra and B. Kalluraya, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2010, **66**(12), o3252.
- 19 D. Grossie, L. Harrison and K. Turnbull, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2014, **70**, o1165–o1166.
- 20 G. B. Riddle, D. A. Grossie and K. Turnbull, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2004, **60**, 2003–2004.
- 21 D. A. Grossie, K. Turnbull, S. Felix-Balderrama and S. Raghavapuram, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2009, **65**, o554–o555.
- 22 C. Favre and F. Friscourt, *Org. Lett.*, 2018, **20**, 4213–4217.
- 23 H. Hope and W. E. Thiessen, *Acta Crystallogr., Sect. B*, 1969, **25**, 1237–1247.
- 24 H. K. Fun, T. S. Chia, Nithinchandra, B. Kalluraya and S. Shetty, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2012, **68**(7), o2103.
- 25 Y. Dürüst, E. Yıldız, H. Karakuş and B. M. Kariuki, *Synth. Commun.*, 2017, **47**, 660–670.
- 26 T. J. King, P. N. Preston, J. S. Suffolk and K. Turnbull, *J. Chem. Soc., Perkin Trans. 2*, 1979, 1751.
- 27 I. S. Khazhieva, P. M. Demkin, J. INein, T. V. Glukhareva and Y. Y. Morzherin, *CCDC 1030491: Experimental Crystal Structure Determination*, 2016.
- 28 I. A. Cherepanov, A. S. Samarskaya, I. A. Godovikov, K. A. Lyssenko, A. A. Pankratova and V. N. Kalinin, *Tetrahedron Lett.*, 2018, **59**, 727–729.
- 29 M. Riomet, K. Porte, L. Madegard, P. Thuéry, D. Audisio and F. Taran, *Org. Lett.*, 2020, **22**, 2403–2408.
- 30 V. N. Kalinin, S. N. Lebedev, I. A. Cherepanov, I. A. Godovikov, K. A. Lyssenko and E. Hey-Hawkins, *Polyhedron*, 2009, **28**, 2411–2417.
- 31 V. A. Budevich, S. V. Voitekhovich, A. V. Zuraev, V. E. Matulis, V. E. Matulis, A. S. Lyakhov, L. S. Ivashkevich and O. A. Ivashkevich, *Beilstein J. Org. Chem.*, 2021, **17**, 385–395.
- 32 P. S. V. Lima, G. H. Weimer, L. P. Oliveira, H. D. S. Souza, G. F. Fiss, H. G. Bonacorso and M. A. P. Martins, *CrystEngComm*, 2023, **25**, 4976–4991.
- 33 D. Cantillo, M. Ávalos, R. Babiano, P. Cintas, J. L. Jiménez, M. E. Light, J. C. Palacios and V. Rodríguez, *Org. Biomol. Chem.*, 2010, **8**, 5367–5374.
- 34 L. B. de Oliveira Freitas, P. Eisenberger and C. M. Cradden, *Organometallics*, 2013, **32**, 6635–6638.
- 35 G. Herrschaft and H. Hartl, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1989, **45**, 1021–1024.

36 H. Ishigami, M. Sumita, Y. Tsunashima, T. Hori and S. Sato, *J. Korean Phys. Soc.*, 2003, **42**, 1237–1239.

37 U. Böhme and M. Gerwig, *CCDC 1874521: Experimental Crystal Structure Determination*, 2018.

38 D. J. Evans and D. L. Hughes, *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.*, 1990, **46**, 1452–1454.

39 J. K. Cockcroft, A. Shamsabadi, H. Wu and A. R. Rennie, *Phys. Chem. Chem. Phys.*, 2019, **21**, 25945–25951.

40 G. R. Clark, G. L. Shaw, P. W. J. Surman, M. J. Taylor and D. Steele, *J. Chem. Soc., Faraday Trans.*, 1994, **90**, 3139.

41 A. B. Burg, *Inorg. Chem.*, 1989, **28**, 1295–1300.

42 R. T. Boeré, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2007, **63**, o3581.

43 A. R. Cowley, A. J. Downs, S. Marchant, V. A. Macrae, R. A. Taylor and S. Parsons, *Organometallics*, 2005, **24**, 5702–5709.

44 K. L. Mears, M. A. Kutzleb, C. R. Stennett, J. C. Fettinger, D. C. Kaseman, P. Yu, P. Vasko and P. P. Power, *Chem. Commun.*, 2022, **58**, 9910–9913.

45 R. Boese, P. Paetzold, A. Tapper and R. Ziembinski, *Chem. Ber.*, 1989, **122**, 1057–1060.

46 J. Kruszewski and T. M. Krygowski, *Tetrahedron Lett.*, 1972, **13**, 3839–3842.

47 T. M. Krygowski, H. Szatylowicz, O. A. Stasyuk, J. Dominikowska and M. Palusiak, *Chem. Rev.*, 2014, **114**, 6383–6422.

48 K. K. Zborowski, I. Alkorta, J. Elguero and L. M. Proniewicz, *Struct. Chem.*, 2012, **23**, 595–600.

49 A. Huber, J. Dubbert, T. D. Scherz and J. Voskuhl, *Chem. – Eur. J.*, 2023, **29**, 1294.

50 C. W. Bird, *Tetrahedron*, 1985, **41**, 1409–1414.

51 Y. I. Nein and Y. Y. Morzherin, *Russ. Chem. Bull.*, 2012, **61**, 1111–1116.