ABSTRACT: Hydroxides of superalkalis (particularly, K- and Na-related species) are shown for the first time to function as superbases. A new small series of hydroxides (XM_{n+1}OH) is designed based on superalkali species (XM_{n+1}) where M (K and Na) is alkali metal atoms, n is the maximal formal valence of the central atom X (F, O, and N), and n ≥ 1. To probe whether such fascinating polynuclear superalkali hydroxides (SAHs), especially the K- and Na-associated moieties are as basic as the representative alkali metal hydroxides (KOH, NaOH, and LiOH) as well as similar Li-based SAHs, a comprehensive computational exploration (in the gas phase) has been reported using the framework of an ab initio method. The ab initio calculations reveal that both the K- and Na-related SAHs consisting of larger gas-phase proton affinity (PA) and gas-phase basicity (GB) values demonstrate stronger basic character compared to the LiOH and Li-based SAHs. However, the available SAHs act as strong bases as well as superbases; among the proposed K- and Na-based SAHs, remarkably, the OK3OH moiety having the highest PA (1168.4 kJ/mol) and GB (1146.9 kJ/mol) values shows the evidence of the strongest basicity (i.e., superbase/hyperbase), which exceed enough (ΔPA: 142.1 kJ/mol and ΔGB: 146.9 kJ/mol) the IUPAC-defined superbasicity threshold values (PA: 1026.3 kJ/mol and GB: 1000 kJ/mol) of 1,8-bis(dimethylamino)naphthalene (DMAN). Furthermore, theoretical signatures have been predicted via the electronic structure calculation approach in probing the dissociation energy, ionization potential, electron affinity, HOMO−LUMO gap, and chemical hardness as well as the NCI plot and QTAIM tools are used for the bonding feature analysis and such parameters are well linked with the basicity analyzing parameters. In this study, the ab initio-based computational experiments provide some new insights into the basicity features and understanding of the structural and electronic features of a small series of designed K- and Na-related SAHs. Design and synthesis of such theoretically examined SAHs may pave alternative routes for the experimentally rewarding applications.

INTRODUCTION

Molecules having low ionization potentials (IPs) may play a crucial role in chemistry, and alkali metal atoms are quite important because of their very small IPs. In particular, superalkali species show remarkable features due to their lower IPs compared to those (5.39−3.89 eV) of the alkali metal atoms.1 Thus far, examination of superalkali species involves from small to medium size mono-, bi-, and polynuclear species and then unique superalkali. Also, superalkali with outstanding reducing properties have accomplished noteworthy progress earlier and have attracted more and more attention as well as great importance in chemistry. Such success lead to vast construction of new moieties as superalkali. Recently, remarkable studies have been reported regarding the potential utilization of superalkali but the practical application of such species yet persists as a challenge. Studies on the scope of superalkali (especially, synthesis of the nonmetallic and polynuclear superalkali cations associated with diverse functional groups) and superhalogens have greatly expanded in approximately the past 15 years.2−5 Along with such progress, molecular modeling and designing of novel superbases from their simple building blocks are quite promising and are active areas of research and development; however, suitable selection of such model systems for a rational design and synthesis is still to be achieved by experimentalists. By acquiring more pieces of information as well as new insights into the conspicuous geometrical and electronic features of such systems, various theoretical and experimental examinations have been carried out.4−8 Studies on the building blocks of super salts and their related systems like encapsulated superalkali inside fullerene can also be seen in the literature.8−10

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A variety of experiments on molecular species like XLi₂ (X = F, Cl, Br, and I)\textsuperscript{11–13} OM₃ (M = Li, Na, and K)\textsuperscript{14–16} NLi₄, BLi₄, \((n = 1–7)\),\textsuperscript{31} etc. has been reported. As alkali metal and alkali earth metal oxides and their hydroxides (in the gaseous state) are reported to be the strongest bases,\textsuperscript{3} hitherto, a large range of inorganic as well as ionic and covalent organic bases can be viewed, including the bases having high proton affinities (so-called superbases).\textsuperscript{19–22} Alder’s 1,8-bis(dimethylamino)-naphthalene (DMAN) compound (in the gas phase) with proton enthalpy equal to or greater than 1026.3 kJ/mol has been referred a superbase as per recommendation by the International Union of Pure and Applied Chemistry (IUPAC) definition.\textsuperscript{20}

Numerous efforts have been devoted to obtaining experimental findings for synthesizing superalkalis and their related compounds, which have been described along with the synthesis of superhalogens/hyperhalogens.\textsuperscript{12,23–27} For exploring the potential-reducing capabilities of such type of innovative species and synthesis of various charge-transfer (CT) salts, numerous attempts have been made incessantly.\textsuperscript{28} Theoretical characterization of a new type of superhalogen-(super)alkali compound BF₄\textsuperscript{−}M (M = Li, FLi₂, OLi₃, and NLi₄) has also been reported, which revealed the nonlinear optical properties of such superatom compounds.\textsuperscript{29}

Despite the abovementioned achievements, there is still room for improvement and many attempts are still needed to design and characterize methods for acquiring innovative superalkali moieties to enrich the superalkali family. To the best of our knowledge, designing and theoretical characterization (in terms of capability to function as superbases/hyperbases) of \(\text{XM}_{n+1}\text{OH}\) (where \(M = K, N\)) have not been explored previously. In the \(\text{XM}_{n+1}\) segment of the \(\text{XM}_{n+1}\text{OH}\) species, “\(n\)” refers to the number of valence electrons needed to complete the electronic shell of the electronnegative X atom (F, O, and N), and hence, one extra alkali metal atom (M) (\(M = K, N\), and Li) will provide an extra valence electron for the electronic shell of the X atom. Afterward, a high tendency to drop one valence electron is demonstrated by the \(\text{XM}_{n+1}\) molecular system for constructing a very stable cation having positive charges spread over all the “\(n + 1\)” alkali atoms (M). The goal of research work is twofold: first, a small series of hydroxide of superalkalis especially K- and Na-based SAHs has mainly been considered and efforts have been done in obtaining new insights into the absolute gas-phase proton affinity (PA) and intrinsic gas-phase basicity (GB) of these superbases and then comparing with the LiOH and Li-related SAHs. Second, the calculated PA and GB parameters are well linked to some very important and useful analyzed electronic feature-based outcomes. It is contemplated the possibility that the superalkalis, as well as related species, may be brought to play as the core for synthesizing novel superhalogen—superalkali compounds and offering fascinating electronic features.\textsuperscript{27}

In this study, because of the importance of cation/molecule interactions in chemical and biochemical processes, a new series of novel and strong bases \(\text{XM}_{n+1}\text{OH}\) SAHs (such as FK₂OH, OK₂OH, NK₂OH, FN₂OH, ONa₂OH, and NNa₂OH) have been tentatively designed and examined, including the corresponding fundamental bases KOH and NaOH along with LiOH and its allied SAHs acting as strong bases and superbases. The foremost advantage of designing and characterizing theoretically the K- and Na-related SAHs is that these species show their stronger basicity behavior (all act as superbases) than the earlier reported LiOH as well as its related SAHs.\textsuperscript{30} Among all the \(\text{XM}_{n+1}\text{OH}\) species, the K-related SAHs, especially, the \(\text{OK}_3\text{OH}\) moiety consisting of the highest PA and GB values, manifest the strongest basic character. The draught structures of the K- and Na-based moieties show a clear signature of their higher basic character when such species are constructed from a higher number of alkali metals (i.e., superatoms) and hydroxides (OH group). Moreover, attempts have been put forward to obtain some profound and useful insights into the bonding and nonbonding features with the deployment of the noncovalent interaction (NCI) plot and Bader’s quantum theory of atoms in molecules (QTAIM) tools. Several reports on the application of the NCI plot and QTAIM-based topological parameters can be seen in the literature.\textsuperscript{31–34}

**THEORY AND COMPUTATIONS**

The selection of a quantum chemical calculation (QCC) approach that precisely portrays the basicity-related features is an intricate challenge. The choice of an \(\text{ab initio}\) method for bonding between the alkali metal atom (\(\text{XM}_{n+1}\)) and hydroxyl (OH) group of the K- and Na-related SAHs is based on the fact that for the alkali metal and alkaline earth metal oxides and hydroxides, \(\text{ab initio}\) calculations suitably predict the energetic features and also the structural/geometrical outcomes may be in uniformly close agreement with the available experimental data.\textsuperscript{35} The QCC-based optimized (equilibrium) geometries and vibrational harmonic frequencies have been determined by employing the second-order Møller—Plesset (MP2) perturbation theory\textsuperscript{36} in conjunction with a standard split valence basis set including diffuse and polarization functions and the 6-311+\(\text{+G}(d, p)\) basis set using the Gaussian 09\textsuperscript{37} electronic structure calculation program.

A hypothetical gas-phase protonation reaction has been expressed in eq 1 for evaluating the basicity of diverse \(\text{XM}_{n+1}\text{OH}\) hydroxides of K-, Na-, and Li-related species.

\[
\text{XM}_{n+1}\text{OH} + H^+ \rightarrow \text{XM}_{n+1}\text{OH}_2^+
\]  

(1)

where \(\text{XM}_{n+1}\) are FK₂, OK₂, NK₂, FN₂, ONa₂, NNa₂, FLi₂, OLi₂, and NLi₄.

The equation given below (eq 2) describes the accurate gas-phase PA and intrinsic GB of a base (B) (here defined as B = \(\text{XM}_{n+1}\text{OH}\)) and understandably the gas-phase basicity of a neutral base (B) is equal to the acidity of its conjugate acid (BH\(^+\)) (i.e., BH\(^+\) = \(\text{XM}_{n+1}\text{OH}_2^+\)). It should be noted that high values of PA and GB parameters signify higher basicity. For analyzing two very important and useful parameters PA and GB of the designed K- and Na-based SAHs (\(\text{XM}_{n+1}\text{OH}\)) and their protonated forms (\(\text{XM}_{n+1}\text{OH}_2^+\)), the enthalpy (\(\Delta H\)) and Gibbs free energy (\(\Delta G\)) of the given below reaction have been determined

\[
\text{BH}^+(g) \rightleftharpoons B(g) + H^+(g) \quad \text{PA} = \Delta H \quad \text{and} \quad \text{GB} = \Delta G
\]  

(2)

The enthalpy (\(\Delta H\)) and Gibbs free energy (\(\Delta G\)) for the proton (H\(^+\)) have been acquired from the literature.\textsuperscript{38} It is also worth mentioning that the following equation (eq 3) was employed for analyzing the energy of ionic dissociation of the \(\text{XM}_{n+1}\text{OH}\) species into their constituting \(\text{XM}_{n+1}^+\) and OH\(^-\) moieties.

\[
\Delta E = E(\text{XM}_{n+1}^+) + E(\text{OH}^-) - E(\text{XM}_{n+1}\text{OH})
\]  

(3)

where the \(E(\ldots)\) values refer to the total electronic energy of the corresponding species with the zero-point correction.
For the chemical bond study, the electron density (ED)-based chemical bonding (including the weakest bonding as well as nonbonding interatomic interactions) and structural feature analyses of such species have been explored using the wavefunction are conducted with the deployment of Bader’s powerful QTAIM tool. The energetic and topological features in the bond critical point (BCP) are utilized for exploring the type, nature, and strength of the bonding and nonbonding interactions (NBIs)/NCIs. The ED distribution function, \( \rho(r) \), and the wavefunction-based Laplacian, \( \nabla^2 \rho(r) \), can be analyzed for the equilibrium geometry of the probing species by the QTAIM approach with the help of the AIMAll package. The natural bond orbital (NBO) analysis was also employed in evaluating the natural atomic charges wherever needed.

## RESULTS AND DISCUSSION

### XM\(_{n+1}\) Species

We considered the optimized structures and properties of the building blocks (i.e., referred as the superalkalis: XM\(_{n+1}\)) as the starting materials. The optimized geometries of the XM\(_{n+1}\) species are shown in Figure 1 along with some chosen structural parameters particularly, the bond lengths. For example, Figure 1 presents the optimized structures of the neutral K- and Na-related superalkali neutral species. The bond lengths (black color in bold) and IPs (red color in bold) are given in Å and eV, respectively.

### Table 1. Bond Lengths and IPs of the Neutral Superalkali XM\(_{n+1}\) (FK\(_2\), OK\(_3\), NK\(_4\), FNa\(_2\), ONa\(_3\), NNa\(_4\), FLi\(_2\), OLi\(_3\), and NLi\(_4\)) Species Using the MP2/6-311++G(d, p) Level of Theory along with the Natural Charges on the Atoms

| species | symmetry | geometry | bond length (Å) | IP (eV) | natural charge (e) |
|---------|----------|----------|----------------|--------|-------------------|
| K       | -        | -        | -              | -      | -                 |
| FK\(_2\) | C\(_{2v}\) | bent (V shape) | 2.348 (K−F) | 2.934 | F (−0.979e) K (0.49e) |
| OK\(_3\) | D\(_{3h}\) | trigonal planar | 2.367 (K−O) | 2.707 | O (−1.908e) K (0.636e) |
| NK\(_4\) | T\(_d\) (TS) | tetrahedral | 2.686 (K−N) | 1.368 | N (−2.679e) K (0.671e) |
| Na      | -        | -        | -              | -      | -                 |
| FNa\(_2\) | C\(_{2v}\) | bent (V shape) | 2.114 (Na−F) | 3.455 | F (−0.978e) Na (0.489e) |
| ONa\(_3\) | D\(_{3h}\) | trigonal planar | 2.096 (Na−O) | 3.174 | O (−1.915e) Na (0.638e) |
| NNa\(_4\) | T\(_d\) | tetrahedral | 2.173 (Na−N) | 1.822 | N (−1.805e) Na (0.451e) |
| Li      | -        | -        | -              | -      | -                 |
| FLi\(_2\) | C\(_{2v}\) | bent (V shape) | 1.704 (Li−F) | 3.851 | F (−0.941e) Li (0.471e) |
| OLi\(_3\) | D\(_{3h}\) | trigonal planar | 1.706 (Li−O) | 3.463 | O (−1.823e) Li (0.607e) |
| NLi\(_4\) | T\(_d\) | tetrahedral | 1.765 (Li−N) | 3.233 | N (−2.397e) Li (0.599e) |

\(^{a}MP2/6-31+G(d, p).\)
NK₄, FNa₂, ONa₃, and NNa₄) where the equilibrium structures of the neutral Li-related superalkalis XLiₙ₊₁ (FLi₂, OLi₃, and NLi₄) have also been displayed for the sake of convenience and comparison purposes. One can perceive that the equilibrium/optimized structures of the F-, O-, and N-substituted superalkalis are in bent (C₂ᵥ), trigonal planar (D₃h), and tetrahedral (Td) shapes, respectively. Moreover, the neutral superalkalis (FLi₂, FNa₂, and FK₂) and their particular cations (FLi₂⁺, FNa₂⁺, and FK₂⁺) are in bent and linear shapes, respectively.

To analyze the protonation behavior/capability of the K- and Na-related species for establishing the superalkali species, the IPs have been analyzed by evaluating the difference between the energies of the ground states of the neutral superalkali atoms (i.e., superalkalis) and their respective cations. The bond lengths and IPs of all the neutral superalkali species (XMₙ₊₁ = FK₂, OK₃, and NK₄) along with the natural charges on the atoms are listed in Table 1. The expected order of bond lengths of the F-, O-, and N-related species is FK₂ (2.348) > FNa₂ (2.114) > FLi₂ (1.704); OK₃ (2.367) > ONa₃ (2.357); NK₄ (2.357)
(2.096) > OLi₃ (1.706); and NK₄ (2.521) > NNa₄ (2.173) > NLi₄ (1.765) (in Å), respectively, which clearly seem to be dependent on the size of alkali metals (K > Na > Li) and the outcomes of the NX₄-related species are in good agreement with the previous studies.⁴⁴ It is interesting to note that in the case of NK₄ superalkali species, using the MP2/6-31+G(d,p) and MP2/6-311++G(d,p) methods, it was found that the cation (NK₄⁺) species shows a Tₐ structure having a minima in the potential energy surface (PES), whereas in terms of symmetry, the former method gives a Tₐ symmetry (having the minima in the PES) and the latter one provides a C₂ᵥ symmetry (showing a transition state (TS) in the PES) for the neutral NK₄ moiety while both approaches give a Tₐ symmetry structure for the same. One of the excellent features of all nine superalkalis is that all of have lower IP values (NK₄ (1.368 eV) < NNa₄ (1.822 eV) < NLi₄ (3.233 eV); FK₂ (2.934 eV) < FNa₂ (3.455 eV) < FLi₂ (3.851 eV); and OK₃ (2.707 eV) < ONa₃ (3.174 eV) < OLi₃ (3.463 eV)) compared to the earlier highlighted respective alkali metal atoms (K, Na, and Li) with IPs in the range of 5.39−3.89 eV and such outcomes are less adequate to ascertain their superalkali potentials.¹⁴ The analyzed natural charges (Q) on the XMₙ₊⁺ species, for instance, the charges on the F and M atoms (Qₓ, Qₓ⁺) of the FM₂ neutral species, FK₂ (−0.979, 0.49) > FNa₂ (−0.978, 0.489) > FLi₂ (−0.941, 0.471), shown in Table 1 also favor the smooth construction of the superalkali-based compounds.

**KOH and XKn⁺OH Species.** As superbases are of theoretical interest and potentially important in organic synthesis,⁴⁵,⁴⁶ a new series of fascinating and interesting K- and Na-related hydroxides of the superalkalis has been designed and computational experiments were performed for the theoretical characterization. A total of eight hydroxide species has been considered with two sets: set 1 consisting of K-based superalkalis (see Figure 2) and set 2 consisting of Na-based
superalkalis (see Figure 3) and each set having four hydroxides, and a third set with the Li-related species (see Figure 4) also consists of four similar hydroxides. For example, the first set of the K-related \((\text{XK}_{n+1}\text{OH})\) SAH optimized/equilibrium structures has three species \(\text{FK}_2\text{OH}, \text{OK}_3\text{OH}, \text{NK}_4\text{OH}\) including the fundamental base \(\text{KOH}\). Various geometries were scanned in finding out the appropriate K- and Na-related SAHs along with the Li-related species \((\text{XLi}_{n+1}\text{OH}: \text{FLi}_2\text{OH}, \text{OLi}_3\text{OH}, \text{NLi}_4\text{OH})\). As shown in Figure 2 \((\text{XK}_{n+1}\text{OH})\) and Figure 3 \((\text{XNa}_{n+1}\text{OH})\), both sets of the species have \(C_{2v}\) symmetry structures and the hydroxyl (−OH) group involved therein is connected to the superalkalis through two alkali metal atoms (i.e., the OH group is linked to two potassium atoms in the \(\text{XK}_{n+1}\text{OH}\) case forming a four-membered ring) corresponding to the minimum energy geometries.

As shown in Figure 2, for the optimized parameters of a linear KOH equilibrium \((C_{\infty v}\) symmetry) structure, the O−K and O−H bond lengths are 2.244 Å and 0.957 Å, respectively, which are close to their corresponding experimental outcomes \((\text{O−K: 2.212 Å and O−H: 0.912 Å})\). Two structures of the \(\text{XK}_{n+1}\text{OH}\) \((\text{FK}_2\text{OH} \text{and O}_3\text{OH})\) have been observed to be planar except the \(\text{NK}_4\text{OH}\) in which the \(\text{NK}_2\) segment \((\text{i.e., K−N−K having the N−K bond length as } 2.691 \text{ Å in } V \text{ shape})\) lying at right side from the center of the ring \((\text{terminal NK2 existing at right side})\) is perpendicular to the H−O<K2 fragment \((\text{i.e., O−K: 2.446 Å and O−H: 0.959 Å})\) located on the left side from the center of the ring.

From Figure 2, it can be seen that the X−K bond lengths associated with the cyclic form of the \(\text{XK}_{n+1}\text{OH}\) SAHs \((\text{F−K in } \text{FK}_2\text{OH: 2.396 Å, O−K in } \text{OK}_3\text{OH: 2.464 Å, and N−K in } \text{NK}_4\text{OH: 2.788 Å})\) are higher than those in the respective superalkalis (see Figure 1) \((\text{F−K in } \text{FK}_2: 2.348 Å, \text{O−K in } \text{OK}_3: 2.367 Å, \text{and N−K in } \text{NK}_4: 2.521 Å)\) as well as KOH \((\text{O−K: 2.244 Å})\). Such findings are possible because of the CT phenomenon occurring in different K atoms of the SAHs to the hydroxyl (OH) group and also the electronegative atoms \((\text{i.e., F, O, and N})\) involved therein. Using the NBO approach, the

![Figure 4](https://pubs.acs.org/acsomega/10.1021/acsomega.1c04395)
natural charges ($Q_\text{N}$) on the K atoms of the FK$_2$OH, OK$_2$OH, and NK$_4$OH have been calculated as 0.976e, 0.955e, and 0.903e, respectively (note that the $Q_\text{N}$ on the bridged K atoms are shown here) and such a decreasing trend is due to the enhanced X–K bond lengths with the O–K bond length of KOH (2.244 Å) < FK$_2$OH (F–K: 2.396 Å) < OK$_2$OH (O–K: 2.486 Å) < NK$_4$OH (N–K: 2.788 Å) altering from FK$_2$OH to NK$_4$OH species.

The NCI-reduced density gradient (NCI-RDG) tool has been used for the identification and graphical visualization of the bonding (i.e., covalent) and nonbonding (i.e., ionic and NCIs like H-bonds, and van der Waals (vdW) interactions regions. Color schemes provide comprehensive information about the NCIs. The RDG isosurface (2D scatter plot) shown on the horizontal axis ranges from −0.05 to +0.05. On the horizontal axis (between −0.05 and + 0.05), favorable (bonding and nonbonding) interactions emerge on the left side while unfavorable interactions emerge on the right side, and the extremely weak vdW interactions are close to zero (for instance, see the 2D scatter plot shown in Figure 3 (just above the light blue color horizontal line) for XNa$_{11+}$ species). The colored surfaces of all the SAHs have been shown on the vertical axis (right side) in a red–green–blue color palette via $\Delta \rho(r)$ values ranging from −0.035 to +0.02 au (see Figure 3). The higher-density values ($\Delta \rho(r) < 0$) show stronger attractive interactions, which are clarified by the meticulous depiction of the NCI technique, whereas repulsive interactions can be seen by the very low-density values ($\Delta \rho(r) > 0$). For the sake of convenience and clarity, consider the NK$_4$OH species in which the blue spike (between −0.03 and −0.04 au) in the NCI 2D scatter plot and blue lenticils in the 3D isosurface show strong O–K interactions adjacent to the O–H covalent bond. This structure also shows weak N–K bonding interactions with respect to the O–K interaction indicated by the dark green disc shape (in the 3D isosurface map) in the ring bond and dark green spikes in the 2D scatter plot (between −0.015 and −0.025 au), which appear to confirm its weak metal–nonmetal bonding feature similar to a very strong H-bonding interaction. Moreover, the NCI plot of the NK$_4$OH base shows much stronger N–K bonding interactions located at the terminal N–K bonds (two green–blue color discs in the 3D isosurfaces) compared to those of the ring N–K bonds (two dark green color discs in the 3D isosurfaces), which can clearly be verified by the QTAIM picture (vide infra). A small light green-colored peak (2D scatter plot ranging between 0 and 0.005 au) and a small light green disc lying at the center of the ring (3D isosurface map) can also be seen in Figure 3, which reveal the vdW (extremely weak) interaction between the two electronegative O and N atoms (i.e., a bond path as O⋯N). One can see much weaker N–K bonding interactions (i.e., both ring and terminal bonds) (example of X–K) in NK$_4$OH (a clear picture taken from the QTAIM analysis, see detailed discussion below) compared to the O–K and F–K bonding interactions associated with the OK$_2$OH and FK$_2$OH species, respectively (two blue color discs in the ring bonds in both the moieties).

To attain some profound insights, such kinds of chemical bonding features were also corroborated by the QTAIM tool. As this technique provides a good understanding of the type, nature, and strength of bonding and nonbonding properties, hence, to assure such kinds of characteristics, the QTAIM method has been employed. The sign of the Laplacian of the ED ($\nabla^2 \rho$) at the BCP appeared to show depleted or “closed shell” (ionic and vdW) interactions for the O–K and N–K bond lengths (i.e., for ionic interaction, $\nabla^2 \rho > 0$ and $\rho$ is typically small in the order of $10^{-2}$ atomic unit (au) and $10^{-3}$ au for a vdW interaction) whereas “shared” (covalent) interaction for the OH bond, that is, $\nabla^2 \rho < 0$ and $\rho > 0.1$ au (see Figure 2). For example, the QTAIM topological parameters ($\rho$, $\nabla^2 \rho$, $\rho$) for the O–K and N–K interactions probed in NK$_4$OH are $(0.031, +0.162, −0.031)$ and $(0.0219, +0.078, −0.017)$ (in au), respectively, which fulfill their ionic character ($\nabla^2 \rho > 0$ and $\rho \sim$ order of $10^{-2}$ au), where the O–K ionic interaction is much stronger than the N–K ionic interaction as expected. The above findings have also been highlighted by the QTAIM pictures of the NK$_4$OH species in which the O–K bonds are shown by a solid line (i.e., showing stronger interactions) and N–K bonds are represented by dotted lines (i.e., showing weaker interactions with respect to the O–K bond). In the NK$_4$OH, the NK$_4^+$ moiety possesses a high positive natural charge ($0.991e$); in fact, the hydroxyl (OH) group having a negative natural charge (−0.991e) merely interacts with the NK$_4^+$ moiety and thus show ionic interactions, which were also validated by the QTAIM-based $\rho$ and $\nabla^2 \rho$ topological parameters. Similarly, the OK$_3^+$ moieties (consisting of higher positive natural charges) interact only with the OH groups having higher negative natural charges of −0.973e and −0.979e and thus show an ionic character, which were also confirmed by the QTAIM-based parameters. Moreover, the ($\rho$: 0.3596 au, $\nabla^2 \rho$: −2.22 au, and V: −0.763 au) values of the OH bond involved in the NK$_4$OH species are characteristic of its covalent nature where the $\rho$ value is in the order of $10^{-1}$ au (ρ > 0.1 au) and the $\nabla^2 \rho$ value is less than zero ($\nabla^2 \rho < 0$). This (covalent nature) is also true for all OH bonds existing in all the inspected moieties.

In sequence, the order of the natural charges on the (O, H) atoms of the OH groups of XNa$_{11+}$OH is KOH (−1.413e, 0.428e) > FK$_2$OH (−1.402e, 0.423e) > OK$_2$OH (−1.389e, 0.416e) > NK$_4$OH (−1.388e, 0.397e) in which the charges on the O atom of the latter two (OK$_2$OH and NK$_4$OH) are quite close. Very interestingly, the O–H frequency ($\nu$) for the same species follows the trend as KOH (3961.4 cm$^{-1}$) > NK$_4$OH (3940 cm$^{-1}$) > FK$_2$OH (3934.4 cm$^{-1}$) ~ OK$_2$OH (3934 cm$^{-1}$), which is almost consistent with the QTAIM-based parameters ($\rho$, $\nabla^2 \rho$, V) as KOH (0.3602, −2.331, −0.776) > NK$_4$OH (0.3596, −2.22, −0.763) > FK$_2$OH (0.3577, −2.277, −0.767) > OK$_2$OH (0.3577, −2.266, −0.766) (in au). Moreover, as the QTAIM-analyzed ($\rho$, $\nabla^2 \rho$, V) values are characteristic of bond-strength-measuring parameters, the order of the strength of the ring O–K bonds of the XNa$_{11+}$OH including the KOH is KOH (0.445, 0.275, −0.056) > NK$_4$OH (0.031, +1.612, −0.031) > FK$_2$OH (0.0294, +0.15, −0.029) > OK$_2$OH (0.0278, +0.142, −0.027) (in au) showing ionic character ($\rho$ ~ in the order of $10^{-2}$ au and $\nabla^2 \rho > 0$). Additionally, if we have a look into the X–K bonding feature in the same set of the species, the strengths ($\rho$, $\nabla^2 \rho$, V) of the X–K bonds follow the order OK$_2$OH (F–K: 0.0326, +0.169, −0.035) > NK$_4$OH (O–K: 0.0322, +0.151, −0.032) > OK$_2$OH (N–K: 0.0219, +0.078, −0.017) (in au), which are in accordance with the order of the natural charges ($Q_\text{N}$) on the ring K atoms of XNa$_{11+}$OH species (FK$_2$OH: 0.976e) > (OK$_2$OH: 0.955e) > (NK$_4$OH: 0.903e), which are in the reverse order of the X–K bond lengths of FK$_2$OH (F–K: 2.396 Å) < OK$_2$OH (O–K: 2.486 Å) < NK$_4$OH (N–K: 2.788 Å) as expected. A reciprocal trend of the natural charges ($Q_\text{N}$) on the X atoms for the same moieties is FK$_2$OH ($Q_\text{N}$: −0.973e) <
OKOH (Q_E: −1.89e) < NKOH (Q_E: −3.675e) is hoped where the largest enhanced negative charge on the N atom of NKOH is due to its connection with four electropositive K atoms (having a better electron-donating tendency). Also, the natural charges on the O and F atoms of the OKOH (Q_E: −1.89e) and FKOH (Q_E: −0.973e) species are successively in a decreasing order, which are linked with three and two K atoms, respectively. It is appealing to note that the bond lengths of the hydroxyl group (O−H) for all four K-related species range from 0.957 to 0.959 Å (see Figure 2). In addition, there is a good resemblance between the optimized structures of all SAHs (XNa3OH) (Figure 2) and their respective superalkalis (Figure 1).

**NaOH and XNa3+OH Species.** Now, proceeding from K-related species toward the Na-related moieties, in the case of an optimized linear NaOH fundamental base (see Figure 3), the Na−O and O−H bond lengths have been determined as 1.98 and 0.954 Å, respectively, whose values are close to the experimental values (Na−O: 1.95 Å and O−H: 0.941 Å).\(^2\) As discussed above about the structural information of the XNa3+OH-related bases, the optimized XNa3+OH set of bases also has similar geometrical features (in terms of symmetry and geometry).

Figure 3 shows that the bond lengths in the XNa3+OH bases such as X=Na (F=Na in FNa2OH: 2.157 Å, O=Na in ONa3OH: 2.178 Å, and N=Na in NNa4OH: 2.328 Å) were detected to be larger than those in the corresponding superalkalis (F=Na in FNa2: 2.114 Å, O=Na in ONa3: 2.096 Å, and N=Na in NNa4: 2.173 Å) and also in the NaOH base (O=Na: 1.98 Å). Such findings are due to the CT taking place from the alkali metal (Na atoms) to the OH group as well as the ring electronegative (F, O, and N) atoms. In addition, a decreasing trend of the natural charges (Q_E) on the Na atoms (with an approximate 0.02e difference) in the NaOH (0.984e), FNa2OH (0.967e), ONa3OH (0.947e), and NNa4OH (0.927e) can be seen, which is expected evidently because of increased X−Na bond lengths (including the O−Na bond length of NaOH) proceeding from the NaOH alkali hydroxide to its SAHs (i.e., NaOH (1.98 Å) < FNa2OH (F=Na: 2.157 Å) < ONa3OH (O=Na: 2.178 Å) < NNa4OH (N=Na: 2.328 Å)). One-to-one structural similarity between the equilibrium structures of the XNa3+OH SAHs (Figure 3) and XNa3+OH superalkalis (Figure 1) can also be seen.

Now, having a closer look into the NCI-based 2D scatter plots and 3D isosurface maps of the XNa3+OH species, such graphic visualizations give a similar pattern like that for the above-discussed XKO3+OH-related species, and especially, the N−Na ring bonds (greenish-blue color discs) involved in the NNa4OH base show stronger interactions than the N−K ring bonds (dark green color discs) present in the NKOH moiety. One can see that the extremely weak vdW interactions (green color spikes in the 2D scatter plots ranging between 0 and −0.01 au as shown in Figure 3) in the XNa3+OH-related bases are also validated by their corresponding 3D isosurfaces (see the small lentils at the center of the rings). The QTAIM bond strength-measuring indices (ρ, V_2ρ, V) also confirm that the N−Na bonds of the NNa4OH have higher (0.0227, +0.138, and −0.024 au) values (i.e., showing stronger metal−nonmetal ionic bonding) than the (0.0219, +0.078, and −0.017 au) values of the N−K bonds of the NKOH SAH; however, such N−K bonds are much weaker than those of F−Na (0.0279, +0.2, −0.0363 au) and O−Na (0.0308, +0.207, −0.037 au) of the same system. In order to examine the X−Na (F−Na, O−Na, and N−Na) bonding features present in the XNa3+OH species, the bond strength-quantifying parameters (ρ, V_2ρ) show ionic behavior, which is also supported by their respective local potential energy densities (V_2ρ) (V_O−N: −0.37 au), (V_F−N: −0.36 au), and (V_N−N: −0.24 au), respectively.

The (ρ, V_2ρ, V) values for the O−Na bonds of the NaOH, FNaOH, ONa3OH, and NNa4OH species are (0.0431, +0.38, −0.067), (0.0296, +0.216, −0.0364), (0.0282, +0.204, −0.034), and (0.0294, +0.214, −0.036), respectively, showing ionic character, which follow the QTAIM-based ionic criteria (ρ > 0 in the order of 10−2 au and V_2ρ > 0). By extracting information about the O−Na nonmetal-metal bonding interactions, another nonmetal−metal (X−Na) bonding interactions in all the Na-related species have also been examined by the QTAIM topological parameters, and here, for example, the (ρ, V_2ρ) values (in au) of the N−Na (0.0227, +0.138) bonds in the NNa4OH system also appear to be the characteristic of ionic character (ρ value is in the order of 10−2 au and V_2ρ > 0) having the V value as −0.024 au where the O−Na bonds (indicated by black solid lines) are much stronger than those of the N−Na bond(s) (shown by black dotted lines).\(^3\) This is because of the stronger attractive interaction occurring between the O and Na atoms in the O−Na bonds as compared to those between the N and Na atoms in the N−Na bonds. A noteworthy observation (O−Na versus X−K bonding interactions) can be corroborated from the FNa2OH (the O−Na bond (coupled with an OH covalent bond) is stronger than the F−Na bond) and ONa3OH (the O−Na bond (linked to the OH covalent bond) is weaker than the O−Na bond) species (see Figure 3) in which an opposite trend can be seen which could be due to the electronegativity difference as well as the number of electropositive Na atoms taking part therein. Moreover, the QTAIM parameters (ρ, V_2ρ, V) (in au) for the OH bonds of the Na-related species range from (0.358, −2.319, −0.772) to (0.362, −2.384, −0.788) showing covalent character (ρ > 0.1 au and V_2ρ < 0). The higher natural charges on XNa3+OH and their associated OH groups of the XK3+OH species demonstrate that the FNa2+, ONa3+, and NNa4+ moieties having positive natural charges as 0.967e, 0.947e, and 0.927e, respectively, interact only with the corresponding OH groups consisting of opposite negative natural charges as −0.967e, −0.947e, −0.927e and confirm the existence of ionic interactions, which is also assured by the QTAIM tool.

Using the NBO approach, the trend of the natural charges (Q) on the (O, H) atoms for the Na-related species has been computed as NaOH (−1.419e, 0.436e) > FNa2OH (−1.403e, 0.435e) > ONa3OH (−1.383e, 0.428e) > NNa4OH (−1.369e, 0.425e) where the charge on the H atoms of NaOH and FNa2OH is approximately the same. Moreover, the Na-series systems consisting of higher O−H stretching frequencies as NaOH (4012.5 cm−1) > FNa2OH (3969.8 cm−1) > ONa3OH (3967 cm−1) > NNa4OH (3958 cm−1) as compared to the corresponding set of the K-series bases, and here, the frequencies (ν) trend of the O−H covalent bonds for the XNa3+OH species (including the NaOH base) are slightly from that of the K-related species. The force constants of the respective bonds in the Na-related SAHs also vary as F−Na > O−Na > N−Na. The negative charge on the F atom (Q_F: −0.967e) of FNa2OH is observed to be the smallest because of its bonding with only two electropositive Na atoms; however, in the case of ONa3OH and NNa4OH, the largest negative natural charges on the O (Q_O: −1.903e) and N (Q_N: −2.55e) atoms are analyzed, which could be due to their connection with three and four electropositive
Na atoms, respectively. The O–H stretching frequency findings come up with the same order of the some important and selected QTAIM wavefunction-based parameters ($\rho$, $V^2\rho$, $V$) as NaOH (0.3626, −2.384, −0.788) > FNaOH (0.3596, −2.34, −0.777) > ONaOH (0.3595, −2.326, −0.775) > NNaOH (0.358, −2.319, −0.772) (in au), which shows their covariant character and this feature is assisted by the QTAIM-based covariant criteria ($\rho > 0.1$ au and $V^2\rho < 0$). It is also interesting to observe that the natural charges on the OH-related O atom of the ONaOH (−1.383e) and NNaOH (−1.369e) have been computed to be smaller than those of the OKOH (−1.389e) and NKOH (−1.388) species, which could be because of higher number of K atoms engaged therein (latter two species); however, an opposite trend can be seen in the case of the H atoms taking part therein (ONaOH (0.428e) = KOH (0.428e) > OKOH (0.428e) > ONaOH (0.428e) = NNa4OH (0.425e) > NNa4OH (0.425e) and NNa4OH (0.425e) ~ FKOH (0.423e) > NKOH (0.397e)).

**LiOH and XLi$_{+1}$OH Species.** As discussed above regarding the Na- and K-based SAHs, having a quick look into the Li-related systems (i.e., mainly the NCI plot and QTAIM analysis have been emphasized here), glimpses on the Li-related systems is provided in this section. The bond lengths (Li−O: 1.607 Å and O–H: 0.952 Å) of the optimized linear LiOH base (see Figure 4) are close to the experimental values (Li−O: 1.582 Å and O–H: 0.969 Å). The X−Li bond lengths of the XLi$_{+1}$OH SAHs are larger than those of the corresponding Li-based superalkalis (XLi$_{+1}$: FLi$_2$O, OLi$_2$O, and NLi$_2$O and also the LiOH base. Such alterations seem to be due to the CT process from the engaged Li atoms to the OH group and also to the other electronegative monovalent (F), divalent (O), and trivalent (N) atoms taking part therein. The NCI plots and QTAIM pictures displayed in Figure 4 show that the bonds associated in the LiOH and XLi$_{+1}$OH are stronger than the corresponding bonds involved in both the above discussed XNa$_{+1}$OH as well as XK$_{+1}$OH species.

The NCI-based 2D scatter plots of the Li-associated SAHs clearly illustrate that the blue colored spikes are close to −0.04 showing stronger bonding interactions compared to those in the K- and Na-related fundamental bases and allied SAHs including the LiOH having a blue peak beyond −0.05 whereas in the KOH and NaOH bases, peaks are in between −0.04 and −0.05. The NCI 3D isosurfaces (blue color lentils lying between the O–Li and X−Li ring bonds) also exemplify and authenticate the 2D scatter plots. Stronger vdW interactions with respect to the XK$_{+1}$OH and XNa$_{+1}$OH species can be seen in Figure 4 (2D scatter plots and 3D isosurface maps), which may further be corroborated by both the 2D scatter plots (i.e., spikes lying between −0.01 and −0.02) and 3D isosurface maps (green color lentils located inside the ring). Moreover, such above-highlighted vdW interactions (i.e., see the diagonal connection: O···F@FLi$_2$OH, O···O@OLi$_2$OH, and O···N@N Li$_2$OH shown in the pink and black dotted colors in the 3D isosurface maps and QTAIM pictures, respectively) seen in the XLi$_{+1}$OH species appear to provide extra stability to all the ring-associated moieties, which could be due to their dominating close contacts diagonally and thus demonstrating more attractive interactions compared to other two sets of the SAHs, XK$_{+1}$OH and XNa$_{+1}$OH. In terms of some bond strength-quantifying parameters ($\rho$, $V^2\rho$, $V$), for all the XLi$_{+1}$OH SAHs, the order of the bond strengths of diagonally connected atoms is O···F (0.0118, +0.059, −0.018) > O···O (0.0175, +0.049, −0.016) > O···N (0.016, +0.039, −0.014) (in au), which reveals to be dependent on the number of extra Li atoms (excluding the ring/bridged Li atoms) involved in the species because the presence of extra terminal Li atom enhances the negative charge on the X atom, which may be responsible for more repulsion with the diagonally connected O atom present on opposite side.

In order to probe the order of the bond strength of the OH-associated O−Li bonding interactions, a pattern like LiOH (0.0635, +0.635, −0.115) > FLi$_2$OH (0.0407, +0.399, −0.057) > OLi$_2$OH (0.0388, +0.322, −0.054) > N Li$_2$OH (0.0387, +0.318, −0.053) (in au) has been observed in all four Li-related moiety systems acquired from the QTAIM-based ($\rho$, $V^2\rho$, $V$) parameters having much larger values (i.e., stronger) than the OH-coupled O−M (O−K and O−Na) bonds of the XM$_{+1}$OH species, and also the OH-connected O−Li bonds show stronger ionic character (ρ ~ in the order of 10$^{-3}$ au and V$^2\rho > 0$) than the O−K and O−Na bonds. The above trend is opposite to their respective O−Li bond lengths as LiOH (1.607 Å) < FLi$_2$OH (1.796 Å) < OLi$_2$OH (1.81 Å) < N Li$_2$OH (1.812 Å) as expected. Like the XNa$_{+1}$OH species, the order of the X−Li (X = F, O, and N) bond strength of the XLi$_{+1}$OH SAHs obtained from the ($\rho$, $V^2\rho$, $V$) parameters is (OLiOH (O−Li: 0.0443, +0.344, −0.062)) ≥ (FLi$_2$OH (F−Li: 0.0426, +0.36, −0.061)) > (NLi$_2$OH (N−Li: 0.0361, +0.239, −0.045)) (in au); however, their corresponding bond lengths are O−Li (1.796 Å), F−Li (1.752 Å), and N−Li (1.917 Å), which shows that such properties do not depend only on the bond lengths but also on the natural charges on the constituents. The analyzed natural charges on the bridged Li atoms of the XLi$_{+1}$OH (FLi$_2$OH, OLi$_2$OH, and N Li$_2$OH) are 0.91e, 0.871e, and 0.817e, respectively, along with the LiOH having the highest natural charge (Q$_{Li}$) on the Li atom as 0.96e. Such a consequence is possible due to the increased bond lengths from the LiOH base to the N Li$_2$OH SAH. In addition, having a view into the O−H stretching frequency of the XLi$_{+1}$OH species, a trend as LiOH (4049.8 cm$^{-1}$) > FLi$_2$OH (4016.8 cm$^{-1}$) > OLi$_2$OH (4092.2 cm$^{-1}$) > N Li$_2$OH (3996 cm$^{-1}$) can be seen, which is in close proximity to the order of the QTAIM-analyzed ($\rho$, $V^2\rho$, $V$) parameters as LiOH (0.3647, −2.472, −0.799) > FLi$_2$OH (0.3621, −2.432, −0.79) > OLi$_2$OH (0.3618, −2.409, −0.788) > NLi$_2$OH (0.3606, −2.392, −0.784) (in au). This pattern is considered with the alteration in their analyzed respective O−H bond lengths LiOH (0.952 Å) < FLi$_2$OH (0.954 Å) = OLi$_2$OH (0.954 Å) < N Li$_2$OH (0.955 Å). Moreover, the bond lengths of the OH group of the XLi$_{+1}$OH superbases ranging from 0.952 to 0.955 Å (see Figure 4) show a marginal change (0.003 Å).

**Dissociation Energy ($\Delta E$ and $D$).** As highlighted above, using the NBO approach, the interaction (i.e., type and nature) between the superalkalis (XM$_{+1}$O) and hydroxyl groups (−OH) has been detected to be ionic as in the KOH and NaOH along with the LiOH, which have been examined by looking into the segregations of the XM$_{+1}$O into the superalkali cations (XM$_{+1}$) and OH$^-$ and then the reactions have been compared with those of the KOH, NaOH, and LiOH. From Table 2, it is interesting to see that the ionic detachment/dissociation energies ($\Delta E$) of all the XM$_{+1}$O have been calculated to be smaller than those of their corresponding basic hydroxides (KOH, NaOH, and LiOH). For instance, the order of $\Delta E$ values in (kJ/mol) for the K-based SAHs is KOH (589) > FK$_2$OH (567.9) > N K$_2$OH (554.7) > OK$_2$OH (516). A similar trend can be seen in the case of Na-related species as NaOH (649.2) > FNaOH (645.1) > N NaOH (594) > ONaOH (546.4) (in kJ/mol). Moreover, a slightly different pattern for the Li-related species can be viewed as LiOH (787.4) > FLi$_2$OH (776.3) > OLi$_2$OH (695) > NLi$_2$OH (662) (in kJ/mol). Such trends are
very close to the XM$_{n+1}$ superalkalis having smaller IPs as compared to those of the corresponding K, Na, and Li alkali metal atoms (see Table 1). For example, in the case of K-related species, the IP values of XM$_{n+1}$ (FK$_2$: 2.934 eV, OK$_2$: 2.707 eV, NK$_2$: 1.368 eV) are much lower than that of the K having an IP value of 4.235 eV.

It should be noted that the IP value of the K-related species has been determined to be smaller than those of Na- and Li-related species. By probing the existence of the XM$_{n+1}$OH$_2$ species at least in the gas phase, fragmentation into stable molecules (namely, FK$_2$OH → KOH + KF; OK$_2$OH → KOH + K$_2$O; and NK$_2$OH → KOH + K$_3$N) has been contemplated. The analyzed detachment/dissociation energies (ΔE$_D$) have been found to be greater than zero (see Table 2) where the order of the D$_2$ values for K-related moieties has been found to be NK$_2$OH (164.6) < OK$_2$OH (197.1) < FK$_2$OH (203.2) (in kJ/mol) except in the case of Na-related (NNa$_2$OH (220.7) < FNa$_2$OH (244.6) < ONa$_2$OH (246.2) and Li-related (NLi$_2$OH (254.4) < FLi$_2$OH (260.5) < OLi$_2$OH (269.1)) (in kJ/mol) showing decreased association of the OH group and superalkalis, which can further be assured by the QTAIM-based topological parameters (ρ, V$\rho$, V) of the O–F bonds as FLi$_2$OH$_2$ (0.0157, +0.024, −0.013) → FNa$_2$OH$_2$ (0.0078, +0.027, −0.006) → FK$_2$OH$_2$ (0.0053, +0.024, −0.004) (in au).

### Table 2. Absolute PA, Intrinsic GB, and Dissociation Energies (ΔE and D$_2$) of the KOH, NaOH, and LiOH and XM$_{n+1}$OH Species (XM$_{n+1}$ = FK$_2$, OK$_2$, NK$_2$, FNa$_2$, ONa$_2$, NNa$_2$, FLi$_2$, OLi$_2$, and NLi$_2$)$^{a,b}$

| species | symmetry | PA  | GB  | ΔE  | D$_2$ |
|---------|----------|-----|-----|-----|------|
| KOH     | C$_{av}$ | 1129.2 | (1101.8)$^d$ | 1104.8 | (1075.4)$^d$ | 589 |- |
| FK$_2$OH | C$_1$   | 1126.9 | 1114.3 | 567.9 | 203.2 |
| FK$_2$OH (ring)$^e$ | C$_{2v}$ | 1103 | 1077.9 | - | - |
| OK$_2$OH | C$_{2v}$ | 1168.4 | 1146.9 | 516 | 197.1 |
| NK$_2$OH | C$_{2v}$ | 1134.7 | 1117.2 | 554.7 | 164.6 |
| NaOH    | C$_{av}$ | 1092.6 | (1071.8)$^d$ | 1066.4 | (1044.8)$^d$ | 649.2 |
| FK$_2$OH$_2$ | C$_1$ | 1073.5 | 1056.5 | 645.1 | 244.6 |
| FK$_2$OH$_2$ (ring)$^e$ | C$_{2v}$ | 1034.1 | 1008.4 | - | - |
| ONa$_2$OH | C$_{2v}$ | 1106.2 | 1081.5 | 546.4 | 246.2 |
| NNa$_2$OH | C$_{2v}$ | 1094.9 | 1079.8 | 594 | 220.7 |
| LiOH    | C$_{av}$ | 1001.3 | (1000.1)$^d$ | 975.9 | 787.4 |
| FLi$_2$OH | C$_1$ | 990.3 | 972.4 | 766.3 | 260.5 |
| FLi$_2$OH (ring)$^e$ | C$_{2v}$ | 939.8 | 912.3 | - | - |
| OLi$_2$OH | C$_{2v}$ | 1027.4 | 1001.7 | 695 | 269.1 |
| NLi$_2$OH | C$_{2v}$ | 1047.7 | 1021.5 | 662 | 254.4 |

$^a$PA, GB, and dissociation energies (ΔE and D$_2$) are in kJ/mol and the E$_{exp}$ values are in eV. $^b$(XM$_{n+1}$OH)$_{ring}$ + (H$^+$) → (XM$_{n+1}$OH)$_{ring}$ has been used for the PA and GB calculations. $^c$(XM$_{n+1}$OH)$_{ring}$ + (H$^+$) → (XM$_{n+1}$OH)$_{ring}$ has been used for the PA and GB calculations. $^d$These data are sorted according to the empirical formula as well as according to the evaluated gas basicity.21

Protonated Superalkali Hydroxide (PSAH) Species (XM$_{n+1}$OH$_2^+$) and Their Absolute PA and GB. Figure 5 shows the optimized (equilibrium) structures of PSAHs (XM$_{n+1}$OH$_2^+$) with some important and chosen structural parameters in which all superalkali hydroxides have been protonated by adding one proton therein. Let us have a look into the optimized structures (Figure 5) and the QTAIM pictures (Figure 6) of all protonated species (XM$_{n+1}$OH$_2^+$), it is revealed that ring diagonal NBIs exist (shown by dotted light gray color lines) between the O atom of the OH group and the X (F, O, and N) atoms of the superalkali groups except the two species (OK$_2$OH$_2^+$ and NK$_2$OH$_2^+$), whose strength reduced going from Li- to Na- to K-related species.

For example, in the case of FM$_2$OH$_2^+$ species, one can see that during the protonation on the OH group, the order of the bond lengths of the O–H bonds (decreases) and the O–F bonds (increases) as FLi$_2$OH$_2^+$ (0.967 and 2.757 Å) → FNa$_2$OH$_2^+$ (0.965 and 3.069 Å) → FK$_2$OH$_2^+$ (0.963 and 3.239 Å) showing decreased association of the OH group and superalkalis, which can further be assured by the QTAIM-based topological parameters (ρ, V$\rho$, V) of the O–F bonds as FLi$_2$OH$_2^+$ (0.0157, +0.024, −0.013) → FNa$_2$OH$_2^+$ (0.0078, +0.027, −0.006) → FK$_2$OH$_2^+$ (0.0053, +0.024, −0.004) (in au).

Using the electronic structure calculation method, precise gas-phase PA and useful information on the structure of the base and its conjugate acid are obtained, and theoretically calculated PAs are more accurate than the relative values. In acquiring the PA and GB parameters for the XM$_{n+1}$OH, a hypothetical protonation reaction has been highlighted in the earlier section (Theory and Computations). To vindicate the existing patterns in the basicity features, it is important to examine the alterations in relevant structural parameters and bond lengths induced by protonation. Very interestingly, upon the protonation, the O–M bond lengths (where M = K, Na, and Li) get elongated in all the XM$_{n+1}$OH$_2^+$ species; for instance, in the KOH and KOH$_2^+$, these are 2.244 and 2.629 Å, respectively and the change in O–K (ΔO–K) is 0.385 Å and similarly the change in O–K (ΔO–K) is 0.561 Å in the case going from NK$_2$OH (2.446 Å) to NK$_3$OH$_2^+$ (3.007 Å). Furthermore, lighter species like the Li-related SAHs including the LiOH base undergo even a smaller change from 1.607 Å in LiOH to 1.869 Å in LiOH$_2^+$ (change: 0.262 Å) or from 1.812 Å in NLi$_2$OH to 2.217 Å in NLi$_2$OH$_2^+$ (change: 0.405 Å). Trends for other species are also in accordance to the above-highlighted pattern, which can be verified from the corresponding O–M bond lengths shown in Figures 2–5. These outcomes demonstrate that the enhancement of the O–M bond length is larger as the basicity is higher, and in general, it occurs as a consequence of a complementary ED transfer (i.e., CT).
from the alkali metals to the OH group. For instance, the NBO charge analysis shows that in the FK2OH2⁺ cation, the charge on the incoming proton is only 0.468e, whereas it changes from −1.402e to −0.933e on the O atom assuring strong electron donation. Along with the above alterations, the enhancement in the charges from 0.976e to 0.988e on the K atom and from −0.973e to −0.979e on the F atom can be seen from Table S1 of the Supporting Information (SI) going from nonprotonated SAH (FK2OH) to protonated SAH (FK2OH2⁺). The O–H stretching frequencies of the protonated as well as nonprotonated SAHs are shown in Table S1, which provide supplementary features.

The analyzed PA and GB values for all superalkali bases are tabularized in Table 2. It is worth mentioning that the basicity of newly designed SAHs (viz., the K- and Na-related SAHs) surpasses the experimental absolute PA (1026.3 kJ/mol) and intrinsic GB (1000 kJ/mol) values of the DMAN (the first known organic compound with high basicity), which has been accepted as a threshold for superbases.20 The PA and GB values for the KOH base have been calculated as 1129.2 and 1104.8 kJ/mol, respectively, which are in close agreement with the PA (1101.8 kJ/mol) and GB (1075.4 kJ/mol) values reported by Lias et al.21 One can note that the PA and GB values of the KOH base are much lower than the corresponding values of the OK3OH (PA: 1168.4 kJ/mol and GB: 1146.9 kJ/mol) (i.e., the highest one among all) and NK4OH (PA: 1134.7 kJ/mol and GB: 1117.2 kJ/mol) (i.e., the second highest among all) and these larger values of the K-based SAHs (OK3OH and NK4OH) are indeed largely sufficient to construct these moieties, which may act as superbases.

However, conversely, the PA (1129.2 kJ/mol) and GB (1104.8 kJ/mol) values of KOH are slightly greater than those of FK2OHlinear (PA: 1126.9 kJ/mol) and significantly larger than those of FK2OHring (PA: 1103 kJ/mol and GB: 1077.9 kJ/mol) superalkali bases yet it acts as a strong base. Such a finding could be elucidated by the occurrence of the F atom (χ = 4) with the
Figure 6. QTAIM pictures and topological parameters ($\rho$, $\nabla^2 \rho$, $V$) of the MOH$_2^+$ and XM$_{n+1}$OH$_2^+$ species.
highest electronegativity) in the FK2OH species (EA: 0.111 eV) having a much higher electron affinity than those of the species containing O (χ: 3.5) (EA for OK3OH is 0.08 eV) and N (χ: 3) (EA for NK4OH is 0.016 eV) atoms (see Table 3). It is

Table 3. IP, EA, HOMO—LUMO Gap (Egap), χ, η, and μ Values of LiOH, NaOH, KOH, and XMO1,OH Species (XMO1, = FK3, OK2, NK2, FNa2, ONa3, NNa4, FLi2, OLi3, and NLi4) a

| Species | IP | EA | Egap | χ | η | μ |
|---------|----|----|------|---|---|---|
| KOH     | 8.78 | 0.179 | 8.601 | 4.48 | 4.301 | 7.683 |
| FK3OH   | 9.458 | 0.08 | 9.347 | 4.785 | 4.674 | 1.14 |
| OK2OH   | 5.788 | 0.08 | 5.708 | 2.934 | 2.854 | 8.111 |
| NK2OH   | 3.322 | 0.016 | 3.306 | 1.669 | 1.653 | 0.029 |
| NaOH    | 9.071 | 0.315 | 8.756 | 4.693 | 4.378 | 7.057 |
| FNa2OH  | 10.19 | 0.14 | 10.05 | 5.165 | 5.025 | 1.616 |
| ONa3OH  | 6.598 | 0.149 | 6.449 | 3.374 | 3.225 | 7.172 |
| NNa4OH  | 4.743 | 0.021 | 4.722 | 2.382 | 2.361 | 5.147 |
| LiOH    | 10.23 | 0.127 | 10.103 | 5.179 | 5.052 | 4.783 |
| FLi2OH  | 11.329 | 0.193 | 11.136 | 5.761 | 5.568 | 1.582 |
| OLi3OH  | 8.242 | 0.066 | 8.176 | 4.154 | 4.088 | 5.213 |
| NLi4OH  | 5.638 | 0.005 | 5.633 | 2.822 | 2.817 | 4.962 |

aIP, EA, χ, and η values are in eV and μ is in Debye.

interesting to note that the Fκ1OH linear species have a slightly smaller PA value (11269.2 kJ/mol) than that of the KOH (11293.3 kJ/mol) while the GA (1114.3 kJ/mol) value is greater than that of the fundamental KOH base (1104.8 kJ/mol).

A similar observation regarding the PA and GB values finding can be seen for the XNa4−OH species (protonated) species. Like the OKOH and NKOH SAHs, the basicity parameters of the ONa3OH (PA: 11062.6 kJ/mol and GB: 1081.5 kJ/mol) and NNa4OH (PA: 1094.9 kJ/mol and GB: 1079.8 kJ/mol) moieties are higher than those in the NaOH base (PA: 1092.6 kJ/mol and GB: 1066.4 kJ/mol) as well as the FNa2OH (PA: 1034.1 kJ/mol and GB: 1008.4 kJ/mol) and FNa2OH (PA: 1073.5 kJ/mol and 1056.5 kJ/mol) SAHs. The calculated PA and GB values of the NaOH base are in general agreement with the experimental values (PA: 1071.8 kJ/mol and GB: 1044.8 kJ/mol). As the PA and GB values for all Na-related SAHs including the NaOH go beyond the threshold PA and GB values of the DMAN. 20 These SAHs show superbasicity character. Similar to the K-related (FK2OH) SAHs, this outcome could be explicated by the presence of the most electronegative F atom in the FNa2OH moiety having a much higher EA value (0.14 eV) than that in the NaOH base (EA: 0.021); however, the EA value 0.149 eV in the case of the ONa3OH species.

Moreover, a clear picture of the protonated forms of the Li-related SAHs XLi+1OH2 + (FLi2OH + , OLi3OH + , and NLi4OH + ) can be seen in Figure 5 along with their optimized parameters; however, except for the ONa3OH and OLi3OH moieties, the PA and GB values for all Li-related species which are smaller than the threshold superbasicity of the DMAN compound taken as a reference. One should note that in terms of the PA and GB values, the basicity of the NLi4OH (PA: 10477.7 kJ/mol and GB: 1021.5 kJ/mol) is 21.4 kJ/mol and 21.5 kJ/mol larger than those of the reference species (DAMAN) (PA: 10263.3 kJ/mol and GB: 1000.0 kJ/mol), whereas the OLi3OH (PA: 20127.4 kJ/mol and GB: 1001.7 kJ/mol) has a very close basicity to those of the DMAN species. Based on the above findings, the OLi3OH SAH falls in border line of the superbasicity feature. Remarkably, the FLi2OH-related species (FLi2OH linear: PA: 990.3 kJ/mol and GB: 972.4 kJ/mol and FLi2OH ring: PA: 939.8 kJ/mol and GB: 912.3 kJ/mol) have been found to be weaker bases compared to the LiOH (PA: 1001.3 kJ/mol and GB: 975.9 kJ/mol), OLi3OH, and NLi4OH, which seems to be due to the presence of the F atom (the most electronegative) that is responsible for its higher EA value (0.193 eV) in the FLi2OH among all the Li-related species.

Finally, a relative comparison is carried out among all the K-, Na- and Li-related species. The order of the basicity (PA, GB) for the fundamental bases is KOH (11292.2, 1104.8) > NaOH (10926.2, 1066.4) > LiOH (1001.3, 975.9) (in kJ/mol) in which the former two fundamental bases exceed the basicity capability of the Alder’s DMAN organic base. Afterward, by looking into the basicity nature of the O-substituted SAHs, a similar trend for the basicity parameters (PA, GB) can be seen as OKOH (1168.4, 1146.9) > ONa3OH (11062.2, 1081.5) > OLi3OH (1027.4, 1001.7) (in kJ/mol) in which the PA and GB values of OKOH are the strongest superbasics among their related K- and Na-related species, respectively (noting that OKOH is the strongest superbase among all the moieties), whereas the latter one (OLi3OH) is the second strongest base among its Li-related species. In the case of the family of NM4OH SAHs, the pattern for the order of basicity is NK4OH (11347.4, 1117.2) > NNa4OH (1094.9, 1079.8) > NLi4OH (1047.7, 1021.5) (in kJ/mol) in which the PA and GB values of NNa4OH are approximately (87 kJ/mol, 95.7 kJ/mol) far greater than those of the NLi4OH base, showing its superbasicity nature.

The Fκ2OH-related series also give similar trends like all the above-highlighted ones where the F-substituted K-related linear (PA: 11269.2 and GB: 1114.8) and (ring (PA: 1103 and GB: 1077.9) (in kJ/mol) bases are stronger than the Na-related linear (PA: 1073.5 and GB: 1056.5) and ring (PA: 1034.1 and GB: 1008.1) bases as well as much stronger than the first reported DMAN organic superbase (PA: 1026.3 and GA: 1000), while the F-associated Li-related both linear (PA: 990.3 kJ/mol and GB: 972.4 kJ/mol) and ring (PA: 939.8 kJ/mol and GB: 912.3 kJ/mol) bases are even weaker bases than the DMAN species.

Frontier Molecular Orbital (FMO). An energy parameter (here referred to as Egap) obtained from the difference between the FMOs namely the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) is a very valuable quantity and it plays a vital role in investigating the chemical reactivity feature. Species that are more reactive have a lower Egap value. Some important and useful electronic parameters such as HOMO—LUMO gap (Egap), IP, electron affinity (EA), absolute electronegativity (χ), chemical hardness (η), and dipole moment (μ) are listed in Table 3. In all the cases, the F-related moieties like FKOH (9.347 eV), FNa2OH (10.05 eV), and FLi2OH (11.136 eV) have slightly larger Egap values concerning their corresponding series. For example, FKOH (9.347 eV) has a larger value than those KOH (8.601 eV) and its other related molecules like OKOH (5.708 eV) and NKOH (3.306 eV).

Such outcomes fairly demonstrate that the FK2OH SAH gets notably more stabilized compared to the KOH base, which appears to be because of the CT to the most electronegative F atom. From the abovementioned values, it is clear that the HOMO—LUMO gaps for the KOH base are larger than those of OKOH and NKOH. Therefore, it is remarkable to mention that the outlook of the trend of the Egap values is FKOH (9.347) > KOH (8.601) > OKOH (5.708) > NKOH (3.306) (in eV), while the order of basicity (PA, GB) for all these four species is in almost an opposite fashion as FKOH (1103, 1077.9) < KOH
Figure 7. HOMO of the LiOH, NaOH, KOH (top), and XM$_{n+1}$OH species (XM$_{n+1}$ = FLi$_2$/FNa$_2$/FK$_2$ (below the top), OLi$_3$/ONa$_3$/OK$_3$ (above the bottom), and NLi$_4$/NNa$_4$/NK$_4$ (bottom)).

(1126.9, 1114.3) < NK$_4$OH (1134.7, 1117.2) < OK$_3$OH (1168.4, 1146.9) (in kJ/mol). Hence, a straightforward connection between the chemical reactivity and basicity can be seen which advocates that the moieties show more basic character with a lower $E_{gap}$ value.

Using Koopmans’ approximation, quantification of IP and EA is possible by considering the negative of the energy eigenvalues of the HOMO (−HOMO) and the LUMO (−LUMO) of the FMOs. Moreover, finite-difference approaches provide two very important parameters, $\chi$ and $\eta$.\(^53\) It is noteworthy to mention that the order of the analyzed (IP, $\chi$, $\eta$) values is FK$_2$OH (9.458, 4.785, 4.674) > KOH (8.78, 4.48, 4.301) > OK$_3$OH (5.788, 2.934, 2.854) > NK$_4$OH (3.322, 1.669, 1.653) (in eV), which are consistent with the chemical reactivity ($E_{gap}$: FK$_2$OH (9.347) > KOH (8.601) > OK$_3$OH (5.708) > NK$_4$OH (3.306) (in eV) and also almost with the basicity (PA, GB) FK$_2$OH (1103, 1077.9) < KOH (1126.9, 1114.3) < NK$_4$OH (1134.7, 1117.2) < OK$_3$OH (1168.4, 1146.9) (in kJ/mol) of the XKN$_{n+1}$OH moieties. Almost reciprocal relationships can be seen between the (IP, $\chi$, and $\eta$) and the (PA, GB) values in the protonation reactions.

A similar trend can be observed in the case of Na-related species in which the three electronic parameters (IP, $\chi$, $\eta$) follow the order FNa$_2$OH (10.19, 5.165, 5.025) > NaOH (9.071, 4.693, 4.378) > ONa$_3$OH (6.598, 3.374, 3.225) > NNa$_4$OH (4.743, 2.382, 2.361) (in eV), which is in accordance (connected either in proportional or reciprocal way) with the chemical reactivity ($E_{gap}$): FNa$_2$OH (10.05) > NaOH (8.756) > ONa$_3$OH (6.449) > NNa$_4$OH (4.722) (in eV) and also almost with the basicity (PA, GB) FN$_{n+1}$OH (1073.5, 1056.8) < NaOH (1092.6, 1066.4) < NNa$_4$OH (1094.9, 1079.8) < ONa$_3$OH (1106.2, 1081.5) < (in kJ/mol) of the XN$_{n+1}$OH moieties; however, in the case of Li-related species, a perfect inverse correlation can be noticed for the same. One can observe that smaller $\chi$ values are associated with higher PA and GB values. The molecular dipole moments (DM) ($\mu$) of the XN$_{n+1}$OH are listed in Table 3. The computed DMs of the fundamental bases KOH, NaOH, and LiOH are 7.683, 7.057, and 4.783, respectively (in Debye), which are in close proximity with the experimental findings (viz., DM for the LiOH: 4.754 ± 0.002 Debye).\(^54\) The DMs of the OK$_3$OH (8.111 Debye) and KOH (7.683 Debye) are computed to be larger than those of the FK$_2$OH (1.14 Debye) and NK$_4$OH (0.029), which implies that the former two species are more polarized compared to the latter two and the direction of $\mu$ can be seen from the OH (hydroxyl group) to the superalkalis in all cases of the XM$_{n+1}$OH species.

For achieving some more insights into the electronic features of the XM$_{n+1}$OH moieties, the HOMO 3D isosurface maps are plotted and shown in Figure 7. One can notice that the HOMOs are located over the atomic orbitals of the O atoms of the hydroxyl group (OH) for all the MOH bases (KOH, NaOH, and LiOH) and XM$_{n+1}$OH (where $n$ = 1) superbases keeping in mind that the basicities of FM$_{n+1}$OH (FK$_2$OH, FNa$_2$OH, and FK$_2$OH) are comparable to those of the MOH bases; however, the HOMOs of the XM$_2$OH and XM$_3$OH are mainly spread over in a different way. For example, the larger size of the Xn$_n$ segment of XM$_{n+1}$OH is responsible for a more delocalized electron, which could explain well the larger basicity of the XM$_{n+1}$OH superbases. In a similar way, the large HOMO lobes of the XM$_2$OH species can be seen near their X$_n$ fragments (OL$_{n+1}$, ON$_{n+1}$, and OK$_{n+1}$), which could explain their stronger basicity features.

CONCLUDING REMARKS

In this study, designing and theoretical characterization of a new and novel series of K and Na-based SAHs have been elucidated including some highlights on the Li-related bases. The ab initio method is used to conduct an inclusive computational investigation on the basicity as well as other electronic features of the XM$_{n+1}$OH SAHs. The employed ab initio findings disclose that the K- and Na-related SAHs have been detected as stronger bases than the LiOH and its SAHs. However, in the existing SAH examples of the strong bases and superbases, OK$_3$OH acting as the superbase has been found to be the strongest base (in the gas phase) with the largest PA (1168.4 kJ/mol) and GB (1146.9 kJ/mol) values among all the probed species, which are
much higher the threshold basicity values (PA: 1026.3 kJ/mol and GB: 1000 kJ/mol) of DMAN conferred by the IUPAC. In addition, to obtain more insights into the probed SAHs, the dissociation energy, IP, EA, HOMO–LUMO gap, and η have also been analyzed using the electronic structure calculation approach, which provides a good correlation with the basicity values (PA and GB). The NCI plot and QTAIM tools give significant information about the bonding and nonbonding features involved therein.

The design and synthesis of such inspected SAHs may pave a way for practical availing applications. As further exploration of the real-time application of superalkali-based hydroxides is required yet, the author is looking forward to a practical investigation of our predictions based on computational experiments. Moreover, theoretical exploration of the stability of superalkalis and associated species in solvents like water, ethanol, acetone, etc. would definitely be valuable as efforts still have not been paid much and access for new applications of superalkalis and related species can be created, which would be of great significance.

■ ASSOCIATED CONTENT

![image]

Natural charges on the protonated (XM$_{n+}$OH) and nonprotonated (XM$_{n+}$OH$_2^-$) SAHs as well as the O–H frequencies (Table S1), bond lengths of O–M (M = K, Na, and Li) in the nonprotonated (XM$_{n+}$OH) and protonated (XM$_{n+}$OH$_2^-$) SAHs and changes therein (Table S2), and Cartesian coordinates of all the K-, Na-, and Li-related species as well as complete reference 37 (Gaussian) (PDF)

Accession Codes

NCI plots (2D scatter plot and 3D isosurface maps) acquired from Multiwfn, VMD, and IrfanView tools are publicly available free of charge. Gaussian 09 software has been used for the electronic structure calculations, and GaussView 6.0 was utilized for the optimized structure presentation. The AIMAll tool used in analyzing the topological parameters can be accessed by purchasing AIMAll professional from http://aim.tkgristmill.com.

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Notes

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■ DEDICATION

This paper is dedicated to Professor P. v. R. Schleyer for his remarkable contribution in the field of Computational Chemistry and Professor Kaman Singh, BBAU Lucknow (A Central University), India for his notable contribution in the area of Physical Chemistry.

1026.3 kJ/mol and GB: 1000 kJ/mol) of DMAN conferred by the IUPAC.
