Cation-Mutation Design of Quaternary Nitride Semiconductors Lattice-Matched to GaN

Zeng-Hua Cai,† Prineha Narang,‡ Harry A. Atwater,§ Shiyou Chen,*† Chun-Gang Duan,† Zi-Qiang Zhu,† and Jun-Hao Chu†

†Key Laboratory of Polar Materials and Devices (MOE), East China Normal University, Shanghai 200241, China
‡Thomas J. Watson Laboratory of Applied Physics, California Institute of Technology, Pasadena, California 91125, United States

Supporting Information

ABSTRACT: The search for new direct bandgap, earth-abundant semiconductors for efficient, high-quality optoelectronic devices, as well as photovoltaic and photocatalytic energy conversion has attracted considerable interest. One methodology for the search is to study ternary and multiternary semiconductors with more elements and more flexible properties. Cation mutation such as binary → ternary → quaternary for ZnS → CuGaS2 → Cu2ZnSnS4 and ZnO → LiGaO2 → Li2ZnGeO4 led to a series of new quaternary chalcogenide and oxide semiconductors with wide applications. Similarly, starting with GaN, ternary nitrides such as ZnSnN2 and ZnGeN2 have been designed and synthesized recently. However, quaternary nitride semiconductors have never been reported either theoretically or experimentally. Through a combination of the Materials Genome database with the first-principles calculations, we designed a series of quaternary nitride compounds I−II−III−IV Ge2N4 (I = Cu, Ag, Li, Na, K; III = Al, Ga, In) following the GaN → ZnGeN2 → I−II−III−IV Ge2N4 mutation. Akin to Li2ZnGeO4, these quaternary nitrides crystallize in a wurtzite-derived structure as their ground state. The thermodynamic stability analysis shows that while most of them are not stable with respect to phase separation, there are two key exceptions (i.e., LiAlGe2N4 and LiGaGe2N4), which are stable and can be synthesized without any secondary phases. Interestingly, they are both lattice-matched to GaN and ZnO, and their band gaps are direct and larger than that of GaN, 4.36 and 3.74 eV, respectively. They have valence band edges as low as ZnO and conduction band edges as high as GaN, thereby combining the best of GaN and ZnO in a single material. We predict that flexible and efficient band structure engineering can be achieved through forming GaN/LiAlGe2N4/LiGaGe2N4 heterostructures, which have tremendous potential for ultraviolet optoelectronics.

I. INTRODUCTION

As a wide-band gap semiconductor, GaN has attracted wide attention for its applications in blue optoelectronic, high-power, and high-frequency devices.1,2 To engineer the band gap and band structure of GaN for ultraviolet applications, ternary AlGaN (AlxGa1−x N)3 and quaternary AlGaInN (AlxGa1−x−y InyN)4−7 alloys have been synthesized. GaN/AlGaN and GaN/AlGaInN superlattices are grown for multiquantum-well-based ultraviolet light-emitting diodes (LEDs).8,9 Although this is the most established method to engineer the band structure of GaN-based devices, the composition nonuniformity and the GaN/AlGaInN lattice-mismatch hindered the improvement of device performance.10 (Note that the homogeneous AlGaInN alloys had been successfully fabricated in recent studies.5−7) Cation-mutation is another method for band structure engineering of semiconductors. For example, in ZnS, two Zn2+ cations can be mutated into one Cu+ cation and one Ga3+ cation, and then a ternary semiconductor CuGaS2 can be derived. In CuGaS2, two Ga3+ can be further mutated into one Zn2+ and one Ge4+, and then a quaternary semiconductor Cu2ZnGeS4 can be derived,11,12 as shown in Figure 1. Ge4+ can also be substituted by Sn4+, and S2− anion by Se2−, and then a series of I−II−IV−VI4 semiconductors can be derived. Among them, Cu2ZnSnSe4 have much smaller band

Received: September 9, 2015
Revised: October 31, 2015
Published: November 2, 2015

DOI: 10.1021/acs.chemmater.5b03536
Chem. Mater. 2015, 27, 7757−7764
gaps (1.0−1.5 eV) than ZnS, making them ideal light absorber materials in thin film solar cells.13−22
This idea for band structure engineering was proposed by Pamplin and Goodman as early as the 1950s,18,19 but the concept did not attract intensive attention until recently when new applications have been explored based on these quaternary semiconductors with more flexible properties than binary semiconductors.20−25 Because there are four chemical elements involved, it becomes possible to tune the lattice constants, band gap, and band edge positions more flexibly; for example, if we design quaternary nitride semiconductors derived from GaN, we may manipulate the band gaps (edges) while fixing the lattice constants to avoid a lattice-mismatch between the quaternary nitrides and GaN, so that their respective superlattices can be easier to grow. On the other hand, it is desirable that the quaternary nitrides crystallize in an ordered structure and not phase-separate at low growth temperature, in contrast with the AlGaN and AlGaInN disordered alloys, which tend to phase-separate at low temperature and require a high deposition temperature to ensure the composition uniformity.

Following this cation-mutation idea (Figure 1b), recently ternary ZnGeN2, ZnSnN2, and their alloys ZnSn1−xGe1−xN2 derived from GaN have been designed and also synthesized successfully. Furthermore, their band gaps can be tuned almost linearly from 3.4 to 2.0 eV through changing the Sn/Ge composition.26,27 As shown in Figure 1b, further mutation of the Zn3+ into the group I and III cations is still possible, which will lead to a series of quaternary nitrides I−III−Ge2N4. However, these quaternary nitrides have never been reported, either theoretically or experimentally. This is in contrast with the case in sulfides or oxides where a large amount of quaternary compounds such as Cu2ZnGeS4, Cu2ZnSnS4, Cu2CdSnS4, and Li2ZnGeO4 have been synthesized with various applications.16,28−32 It is currently unknown whether these quaternary nitrides can be synthesized. If these quaternaries can be synthesized and their band structure and lattice constants can be tuned with more freedom, they may replace (or compete with) the AlGaN, InGaN, or AlGaN alloys and form lattice-matched heterostructures with GaN.

In this paper, we show the design of a series of quaternary nitride compounds I−III−Ge2N4 (I = Cu, Ag, Li, Na, K; III = Al, Ga, In) following the GaN → ZnGeN2 → I−III−Ge2N4 mutation, and we predict their crystal structure and thermodynamic stability based on the first-principles calculations. Most of these nitrides are not stable with respect to phase separation, which explains why the quaternary nitrides have never been reported. There are, however, two key exceptions, LiAlGe2N4 and LiGaGe2N4, which are stable and may be synthesized without secondary phases. Interestingly, their lattice constants are well matched to that of GaN, and their band gaps are both larger than that of GaN, with the valence band edge close to that of ZnO and the conduction band edge close to that of GaN. So they could replace (or compete with) the AlGaN alloys and form lattice-matched GaN/LiAlGe2N4/LiGaGe2N4 superlattices or heterostructures, which would be ideal for ultraviolet optoelectronic devices.

II. DESIGN SCHEME AND CALCULATION METHODS
A schematic plot of the cation-mutation design of the quaternary semiconductors is shown in Figure 1. Similar to the mutation ZnS → CuGaS2 → Cu2ZnGeS4 (or Cu2ZnSnS4, AgZnSnS4), which have been well studied, GaN can also be mutated into ternary ZnGeN2 through mutating the Ga3+ (III) cations into the Zn2+ (II) and Ge4+ (IV) cations, and the ZnGeN2 can be further mutated into CuAlGe2N4 through mutating Zn2+ (II) into Cu+ (I) and Al3+ (III). If the group I and III cations are replaced by I = Cu, Ag, Li, Na, K; III = Al, Ga, In, a series of I−III−Ge2N4 (more generally I−III−IV−V4) quaternary nitrides can be derived from the binary GaN.

One major principle in this cation-mutation is that the chemical balance is always maintained, so Ga3+ is replaced by Zn2+ and Ge4+ (their average valence is equal to that of Ga3+), and Zn2+ is replaced by Cu+ and Al3+. This chemical balance should be maintained everywhere in the crystal structure of ternary and quaternary nitrides. In binary GaN, each N3− anion (Ga3+ cation) is coordinated by four Ga3+ cations (N3− anions), so that it is charge-neutral around each N 3− anion (Ga3+ cation). After the mutation, each N3− needs to be coordinated by two Zn2+ and two Ge4+ in ZnGeN2, in order to satisfy the local charge-neutrality condition everywhere in the lattice. Previous studies in the ZnS → CuGaS2 → Cu2ZnGeS4 mutation showed that the ordered structures that satisfy this local charge-neutrality condition have lower energies than other structures, which is also known as the octet rule (when the anions are in the eight-electron full-shell state, the structure is energetically more stable).12 For ZnGeN2, it was also found that the two lowest-energy structures with the smallest primitive cell also satisfy this condition and can be considered as derived from the 16-atom supercell of the GaN wurtzite structure, as shown in Figure 2b,c. For quaternary I−III− Ge2N4 two structures that satisfy the local charge-neutrality condition can be derived from the those of ZnGeN2 in which all the cations are ordered and each N3− anion is coordinated by one group I+, one group III3+, and two Ge4+ cations. They have the space group P63 (wurtzite-kesterite structure of

Figure 1. Schematic plot of the cation-mutation design of quaternary chalcogenide (a) and nitride (b) semiconductors, derived from binary II−VI ZnS and II−V GaN, respectively. Cu or Li can also be replaced by other group I (I = Ag, Na, K) cations and Al or Ga by other group III (III = In) cations, and subsequently, 15 quaternary nitrides can be designed.
III. RESULTS AND DISCUSSION

Crystal Structure. To predict the most stable crystal structures of these I–III–Ge,N$_4$ compounds, we need to build a series of possible structures. Through replacing the atoms in the wurtzite-based GaN and ZnGeN$_2$ compounds, two I–III–Ge,N$_4$ structures can be derived, as shown in Figure 2d,e. Other structures, including the kesterite, stannite, and primitive-mixed-CuAu structures of Cu$_2$ZnSnS$_4$ and the structures of the inorganic compounds with a 1:1:2:4 stoichiometry (can be found in Materials Genome database), have also been considered. The total energy calculations showed that I–III–Ge,N$_4$ prefers to crystallize in the structure with a space group of P$_c$, same as the wurtzite-kesterite structure of Cu$_2$ZnSnS$_4$ (Li$_2$ZnSnO$_4$). The calculated formation energies of these I–III–Ge,N$_4$ compounds in the wurtzite-kesterite (P$_c$) and wurtzite-stannite (Pmn$_2$) structures are listed in Table 1. Obviously, the wurtzite-kesterite structure is at least 100 meV/formula-unit lower in energy than the wurtzite-stannite one, and all other structures have even higher energy than wurtzite-stannite structure, so wurtzite-kesterite is the most stable structure of I–III–Ge,N$_4$ among all the structures that we considered.

Cu$_2$ZnSnS$_4$ and Li$_2$ZnGeO$_3$ and Pmn$_2$ (wurtzite-stannite), as shown in Figure 2d,e, respectively. In the following, we will show that they have lower energies than other structures that we considered and are among the low energy structures found using the global crystal structure optimization method.

With these quaternary nitride semiconductors designed following the cation-mutation, we studied their crystal structure, electronic and optical properties as well as the thermodynamic stability using the first-principles calculations within the density-functional formalism as implemented in the VASP code. For the crystal structure optimization, the exchange–correlation functional that we used is the generalized gradient approximation (GGA) of Perdew–Burke–Ernzerhof (PBE). Meanwhile, the hybrid exchange–correlation functional (HSE) was used to calculate the electronic structure and optical properties, because GGA usually underestimates the band gap for these compounds, whereas the HSE functional predicted the band gaps with better agreement with experimental results for Cu$_2$ZnSnS$_4$, ZnGeN$_2$, and GaN. An energy cutoff of 450 eV was used in all cases, and the 4 × 5 × 5 Monkhorst–Pack k-point meshes for the Brillouin-zone integration of the 16-atom primitive cell. To predict the thermodynamic stability of the quaternary I–III–Ge,N$_4$ with respect to phase-separation into elemental phases or binary or ternary compounds, all the possible elemental phases or binary or ternary compounds related to the group I and III, Ge and N elements are searched in the Materials Genome (Materials Project) database, and their formation energies are collected from the database.

Table 1. Calculated Formation Energies $\Delta H_f$ (in eV/Formula Unit) of I–III–Ge,N$_4$ (I = Cu, Ag, Li, Na, K; II = Al, Ga, In) in Wurtzite-Kesterite (P$_c$) and Wurtzite-Stannite (Pmn$_2$) Structures, and Their Lattice Constants a, b, and c (in Å) in Wurtzite-Kesterite Structure (P$_c$)

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\Delta H_f$ (P$_c$)</th>
<th>$\Delta H_f$ (Pmn$_2$)</th>
<th>a</th>
<th>b</th>
<th>c</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuAlGe$_2$N$_4$</td>
<td>$-2.57$</td>
<td>$-2.46$</td>
<td>6.354</td>
<td>5.437</td>
<td>5.201</td>
</tr>
<tr>
<td>AgAlGe$_2$N$_4$</td>
<td>$-1.80$</td>
<td>$-1.64$</td>
<td>6.523</td>
<td>5.543</td>
<td>5.318</td>
</tr>
<tr>
<td>LiAlGe$_2$N$_4$</td>
<td>$-4.96$</td>
<td>$-4.79$</td>
<td>6.357</td>
<td>5.443</td>
<td>5.110</td>
</tr>
<tr>
<td>NaAlGe$_2$N$_4$</td>
<td>$-3.71$</td>
<td>$-3.42$</td>
<td>6.578</td>
<td>5.563</td>
<td>5.224</td>
</tr>
<tr>
<td>KAlGe$_2$N$_4$</td>
<td>$-2.17$</td>
<td>$-1.75$</td>
<td>6.825</td>
<td>5.635</td>
<td>5.383</td>
</tr>
<tr>
<td>CuGaGe$_2$N$_4$</td>
<td>$-0.68$</td>
<td>$-0.43$</td>
<td>6.402</td>
<td>5.480</td>
<td>5.249</td>
</tr>
<tr>
<td>AgGaGe$_2$N$_4$</td>
<td>0.08</td>
<td>0.39</td>
<td>6.582</td>
<td>5.586</td>
<td>5.365</td>
</tr>
<tr>
<td>LiGaGe$_2$N$_4$</td>
<td>$-3.11$</td>
<td>$-2.85$</td>
<td>6.410</td>
<td>5.474</td>
<td>5.163</td>
</tr>
<tr>
<td>NaGaGe$_2$N$_4$</td>
<td>$-1.89$</td>
<td>$-1.51$</td>
<td>6.644</td>
<td>5.594</td>
<td>5.274</td>
</tr>
<tr>
<td>KGaGe$_2$N$_4$</td>
<td>$-0.46$</td>
<td>0.06</td>
<td>6.923</td>
<td>5.660</td>
<td>5.419</td>
</tr>
<tr>
<td>CuInGe$_2$N$_4$</td>
<td>0.39</td>
<td>0.83</td>
<td>6.687</td>
<td>5.564</td>
<td>5.383</td>
</tr>
<tr>
<td>AgInGe$_2$N$_4$</td>
<td>0.90</td>
<td>1.50</td>
<td>6.871</td>
<td>5.653</td>
<td>5.503</td>
</tr>
<tr>
<td>LiInGe$_2$N$_4$</td>
<td>$-1.98$</td>
<td>$-1.38$</td>
<td>6.709</td>
<td>5.569</td>
<td>5.300</td>
</tr>
<tr>
<td>NaInGe$_2$N$_4$</td>
<td>$-1.04$</td>
<td>$-0.19$</td>
<td>6.945</td>
<td>5.664</td>
<td>5.413</td>
</tr>
<tr>
<td>KInGe$_2$N$_4$</td>
<td>0.08</td>
<td>1.17</td>
<td>7.200</td>
<td>5.734</td>
<td>5.555</td>
</tr>
</tbody>
</table>

*Note that the quaternary wurtzite-derived structure has lower symmetry than the binary wurtzite structure, so it has three lattice constants.

DOI: 10.1021/acs.chemmater.5b03536
they should be stable in a structure similar to that of GaN. Because the calculated energy of the wurtzite-kesterite structure is the lowest, our following study will be focused on this structure. The calculated lattice constants of \( \text{I}_\text{I} - \text{III} - \text{Ge}_2\text{N}_4 \) in this structure are also listed in Table 1, which may be compared with the future X-ray diffraction measurement.

**Thermodynamic Stability.** Although the most stable crystal structure of \( \text{I}_\text{I} - \text{III} - \text{Ge}_2\text{N}_4 \) (\( \text{I} = \text{Cu}, \text{Ag}, \text{Li}, \text{Na}, \text{K}; \text{III} = \text{Al}, \text{Ga}, \text{In} \)) is determined from the calculations, it is still a question whether they can be synthesized experimentally or whether they are thermodynamically stable. As the component elements increase, the synthesis of the quaternary nitrides becomes more difficult, because these elements may also form many binary and ternary phases which can compete with the quaternary phase in the synthesis process and coexist in the final products.11,14,43,44

In order to predict whether these quaternary nitrides can be synthesized in a certain chemical environment, we introduce the chemical potential of the component elements to describe the chemical environment quantitatively (e.g., \( \mu_{\text{Ge}}, \mu_{\text{Li}}, \mu_{\text{Ga}}, \mu_{\text{N}} \) of LiGaGe\(_2\)N\(_4\)). When the chemical potential of an element is equal to zero, the element is so rich in the environment that its pure elemental phases can be formed, e.g., \( \mu_{\text{Ge}} = 0 \) means that the bulk Ge in diamond structure can be formed. When the chemical potential is decreased to negative, the element becomes poorer in the environment (the partial pressure becomes lower if it is in the gas phase).

If the quaternary nitride (LiGaGe\(_2\)N\(_4\) as an example) can be synthesized and is stable, the following thermodynamic equilibrium condition should be reached

\[
2\mu_{\text{Ge}} + \mu_{\text{Li}} + \mu_{\text{Ga}} + 4\mu_{\text{N}} = \Delta H_f(\text{LiGaGe}_2\text{N}_4)
\]  

where \( \Delta H_f(\text{LiGaGe}_2\text{N}_4) \) is the calculated formation energy of LiGaGe\(_2\)N\(_4\) (the energy change of the formation reaction from elemental phases to the quaternary compound), which is listed in Table 1. Meanwhile, to avoid the leftover (or coexistence) of elemental phases, Li, Ga, and Ge bulk and N\(_2\) gas, in the final products, the following conditions should be satisfied

\[
\mu_{\text{Ge}} < 0, \mu_{\text{Li}} < 0, \mu_{\text{Ga}} < 0, \mu_{\text{N}} < 0
\]

To avoid the formation of the impurity phases such as the binary and ternary nitride compounds, Li\(_3\)N, Li\(_2\)Ga\(_3\), GaN, Ge\(_3\)N\(_4\), Li\(_3\)Ga\(_2\)N\(_2\), and Li\(_5\)Ge\(_3\)N\(_4\), which may compete with LiGaGe\(_2\)N\(_4\) during the synthesis process, the following conditions should be satisfied,

\[
3\mu_{\text{Li}} + 3\mu_{\text{N}} < \Delta H_f(\text{Li}_3\text{N})
\]

\[
\mu_{\text{Li}} + 3\mu_{\text{N}} < \Delta H_f(\text{Li}_3\text{Ga}_2\text{N}_2)
\]

\[
\mu_{\text{Ga}} + \mu_{\text{N}} < \Delta H_f(\text{GaN})
\]

\[
3\mu_{\text{Ge}} + 4\mu_{\text{N}} < \Delta H_f(\text{Ge}_3\text{N}_4)
\]

\[
3\mu_{\text{Li}} + \mu_{\text{Ga}} + 2\mu_{\text{N}} < \Delta H_f(\text{Li}_3\text{GaN}_2)
\]

\[
5\mu_{\text{Li}} + \mu_{\text{Ge}} + 3\mu_{\text{N}} < \Delta H_f(\text{Li}_5\text{Ge}_3\text{N}_4)
\]

Besides the nitrides shown here, a series of Li–Ga (Li\(_3\)Ga\(_2\), Li\(_2\)Ga\(_3\), Li\(_3\)Ga\(_3\), Li\(_3\)Ga\(_4\)) and Li–Ge (Li\(_3\)Ge\(_2\), Li\(_3\)Ge\(_3\), Li\(_3\)Ge\(_4\), Li\(_3\)Ge\(_5\), Li\(_3\)Ge\(_6\), Li\(_3\)Ge\(_7\)) compounds have also been considered, and they impose more limits on the chemical potentials of Li, Ge, and Ga \( (\mu_{\text{Li}}, \mu_{\text{Ge}}, \mu_{\text{Ga}}) \). It has been computationally expensive to determine the crystal structure and calculate the formation energies of all these competing compounds of the quaternary nitrides; however, the recent

![Figure 3. Limit to the regions (black region, if it exists) that stabilize LiAlGe\(_2\)N\(_4\) (a), LiGaGe\(_2\)N\(_4\) (b), LiInGe\(_2\)N\(_4\) (c), and AgAlGe\(_2\)N\(_4\) (d) in the chemical potential space. Here only slices with \( \mu_{\text{Ge}} = -0.1 \) eV are plotted.](image-url)
development of the Materials Project database has accelerated this process significantly.\textsuperscript{12} The calculated formation energies of all the binary and ternary compounds are listed in Tables S1 and S2.

If there exists a certain chemical potential ($\mu_{\text{Ge}}$, $\mu_{\text{Ga}}$, $\mu_{\text{Li}}$, $\mu_{\text{N}}$) that can satisfy all these thermodynamic conditions (1–8), a single-phase sample of the quaternary nitride can be synthesized without the coexistence of any impurity phases in the environment corresponding to this chemical potential. Therefore, we can predict whether LiGaGe$_2$N$_4$ can be synthesized by solving the mathematical equation/inequation arrays and determining the range of ($\mu_{\text{Ge}}$, $\mu_{\text{Ga}}$, $\mu_{\text{Li}}$, $\mu_{\text{N}}$) is independent, and we can choose $\mu_{\text{Ge}}$, $\mu_{\text{Ga}}$, $\mu_{\text{Li}}$, $\mu_{\text{N}}$ as the variables ($\mu_{\text{N}}$ can be derived). The solution of these inequalities falls in a region in the three-dimensional ($\mu_{\text{Ge}}$, $\mu_{\text{Ga}}$, $\mu_{\text{Li}}$) space. In Figure 3b, the region in the $\mu_{\text{Ge}} = -0.1$ eV plane is plotted for LiGaGe$_2$N$_4$. The black area shows the region where ($\mu_{\text{Ge}}$, $\mu_{\text{Ga}}$, $\mu_{\text{Li}}$, $\mu_{\text{N}}$) satisfy all the conditions, and the lines are corresponding to the limit imposed by the binary and ternary compounds (e.g., Ge$_3$N$_4$ (red line) and LiGe$_2$N$_3$ (blue line)). The existence of this region indicates that LiGaGe$_2$N$_4$ can be synthesized from the thermodynamic point of view. On the other hand, the region is quite narrow for $\mu_{\text{Ge}}$, indicating that the control of the Ga chemical potential (richness) is very important for synthesizing single-phase and high-quality LiGaGe$_2$N$_4$ samples. The strictest limit results from GaN and LiGe$_2$N$_3$, because the energy cost for the phase separation LiGaGe$_2$N$_4 \rightarrow$ GaN+LiGe$_2$N$_3$ is small.

Similarly for LiAlGe$_2$N$_4$, there exists a region in the chemical potential space that satisfies all the thermodynamic conditions, as shown in Figure 3a, so it is possible to synthesize LiAlGe$_2$N$_4$. The control of Al richness is very important in the synthesis, because the energy cost for the phase separation LiAlGe$_2$N$_4 \rightarrow$ GaN+LiGe$_2$N$_3$ is small too.

In contrast to LiGaGe$_2$N$_4$ and LiAlGe$_2$N$_4$, all other quaternary nitrides (I–III$\rightarrow$Ge$_3$N$_4$ (I = Cu, Ag, Na, K; II = Al, Ga, In) and LiInGe$_2$N$_4$) do not have a region in the chemical potential space that satisfy all the thermodynamic conditions. Five of them have positive formation energies (listed in Table 1), indicating that the elements prefer to stay at the elemental phases, rather than bind in the quaternary lattice, so they can not satisfy the eq (1) and ineq (2). Other quaternary nitrides have negative formation energies relative to the elemental phases, but the limits from the binary and ternary compounds preclude the existence of a black region, as shown in Figure 3a, so it is possible to synthesize LiAlGe$_2$N$_4$.

The disappearance of a stable region in the chemical potential space can also be simply ascribed to that the quaternary nitrides are unstable with respect to some phase-separation reactions. For example, CuGaGe$_2$N$_4$ does not have a stable region due to the constraint imposed by Ge$_3$N$_4$, Cu$_3$N, and GaN. This kind of constraint can be described by the following phase-separation reaction

\[ 3\text{CuGaGe}_2\text{N}_4 \rightarrow 2\text{Ge}_3\text{N}_4 + 3\text{GaN} + \text{Cu}_3\text{N} \quad (9) \]

which is an exothermic reaction because the calculated energy change ($\Delta E$, shown in Table 2) is negative (i.e., this reaction can proceed spontaneously). This causes the disappearance of the stable region of CuGaGe$_2$N$_4$. A series of the phase-separation reactions of the unstable quaternary nitrides, and their energy changes are listed in Table 2.

In summary, among all these quaternary nitrides we designed, only two of them are thermodynamically stable and can be synthesized experimentally without impurity phases. They are LiAlGe$_2$N$_4$ and LiGaGe$_2$N$_4$. Due to the small stable region in the chemical potential spaces, it is necessary to control the richness of the component elements precisely in the synthesis process.

The element abundance of the source materials in Earth’s crust is 17 ppm for Li, 19 ppm for Ga, 82 000 ppm for Al, 1.4 ppm for Ge, and 20 ppm for N, all of which are more abundant than In (0.16 ppm).\textsuperscript{50} A possible strategy for their growth would be Metalorganic Chemical Vapor Deposition (MOCVD), despite that the available gaseous precursors of Li is limited. Alternatively, sintering the precursor elements or binary nitrides into the LiAlGe$_2$N$_4$ (LiGaGe$_2$N$_4$) target and then sputtering into films would be another possible way to make them into thin films, and NH$_3$ or N$_2$ atmosphere should be provided to overcome possible N loss during the sputtering process. Special attention should be paid to substrate temperature or partial pressure during the MOCVD or sputtering process in order to avoid the decomposition of this quaternary compound into binary or ternary side products.

Electronic and Optical Properties of LiAlGe$_2$N$_4$ and LiGaGe$_2$N$_4$. Because LiAlGe$_2$N$_4$ and LiGaGe$_2$N$_4$ are thermodynamically stable, we study their electronic and optical properties to see whether they are suitable for optoelectronic applications. The calculated band structure and density of states (DOS) of LiAlGe$_2$N$_4$ and LiGaGe$_2$N$_4$ are shown in Figure 4. Evidently, both LiAlGe$_2$N$_4$ and LiGaGe$_2$N$_4$ have direct band gaps, with the valence band maximum (VBM) and the conduction band minimum (CBM) both located at $\Gamma$ point. The gap of LiAlGe$_2$N$_4$ is 4.36 eV, larger than that of LiGaGe$_2$N$_4$ (3.74 eV). The direct nature of the band gaps can be considered as derived from GaN, which also has a direct band gap (3.4 eV) at $\Gamma$ point, indicating that the crystal structure similarity (as shown in Figure 2) also leads to the electronic band structure similarity. Compared to binary GaN and ternary ZnGeN$_2$, the quaternary LiAlGe$_2$N$_4$ and LiGaGe$_2$N$_4$ have larger band gaps, which is in contrast with the monotonic band gap decrease in the ZnS $\rightarrow$ CuGaS$_2$ $\rightarrow$ Cu$_2$ZnGeS$_4$ (or Cu$_2$ZnSnS$_4$). This can be understood
The total and partial DOS of LiAlGe2N4 and LiGaGe2N4 are shown in Figure 4. The top of the valence bands of LiGaGe2N4 is composed mainly of the N 2p states, weakly hybridized with the Ge 4p, Ge 3d, Ga 4p, Ga 3d, and Li 2p states, whereas the bottom of the conduction band is composed mainly of the antibonding states of the hybridization between Ge 4s, Ga 4s, and N 2s states. Because the N 2p orbitals are localized and their hybridization with other orbitals are weak in these nitrides, the highest valence band has a small dispersion, in contrast with the large dispersion of the lowest conduction band, which is composed mainly of the more delocalized Ge 4s, Ga 4s, and N 2s orbitals.

The band component of LiAlGe2N4 is similar to that of LiGaGe2N4, but the top of the valence bands has only N 2p, Ge 4p, Ge 3d, Al 3p, and Li 2p component, and the bottom of the conduction band has only Ge 4s and N 2s component. Therefore, both LiAlGe2N4 and LiGaGe2N4 can be considered as semiconductors with typical s−p band gaps, similar to GaN or other III−V semiconductors with s−p band gaps. The similar band component indicates that continuous and flexible band structure engineering may be possible through forming the heterostructures between the quaternary nitrides and GaN.

Because the valence bands of both compounds have a high density of states (peak) just below the VBM level, we can understand the band gap increases of the quaternary compounds. Comparing the CBM states of GaN and LiGaGe2N4, Ge 4s level is lower than Ga 4s level, which shifts the CBM level down, but the Li 2s level is much higher than Ga 4s level, which shifts the CBM level up, so the two contributions canceled each other. Because the Ge:Li ratio is 2:1, the final result is that the CBM level of LiGaGe2N4 is slightly shifted down, 0.18 eV lower than that of GaN (see Figure 5). Because Al 3s level is also higher than Ga 4s level, the CBM level of LiAlGe2N4 is slightly shifted up, 0.21 eV higher than that of GaN. Considering both the shifts of VBM and CBM levels, we can understand the band gap increases of the quaternary compounds.

Because the valence bands of GaN and LiGaGe2N4, Ge 4s level is lower than Ga 4p level,12 which shifts the CBM level down, but the Li 2s level is much higher than Ga 4s level, which shifts the CBM level up, so the two contributions canceled each other. Because the Ge:Li ratio is 2:1, the final result is that the CBM level of LiGaGe2N4 is slightly shifted down, 0.18 eV lower than that of GaN (see Figure 5). Because Al 3s level is also higher than Ga 4s level, the CBM level of LiAlGe2N4 is slightly shifted up, 0.21 eV higher than that of GaN. Considering both the shifts of VBM and CBM levels, we can understand the band gap increases of the quaternary compounds.

Because the band gaps of GaN and LiGaGe2N4 are both direct, a strong absorption of the UV light is expected, which is confirmed by our calculated absorption spectra and dielectric constants in Figure 6. The absorption coefficient increases rapidly to the order of 10⁵ cm⁻¹ as the photon energy is above the direct band gaps, indicating that the optical transition between the s−p band gaps is very strong. On the other hand, the valence bands of both compounds have a high density of states (peak) just below the VBM level, which also
contributes to the rapid increase of the absorption coefficient. The strong absorption of the UV light makes these quaternary nitrides very competitive for the UV light detector.

As shown in Figure 5, these quaternary nitrides have their valence band edge levels as low as that of ZnO (the VBM level of LiAlGe₂N₄ is even lower than that of ZnO), and their CBM levels can also be shifted above or below that of GaN, indicating that the element mutation provides an efficient method for manipulating the band edge positions. Furthermore, the lattice constants of LiGaGe₂N₄ and LiAlGe₂N₄ are well matched to GaN and ZnO, with a lattice mismatch less than 3%, as shown in Table 1 (the lattice constants of GaN and ZnO in the same supercell as that in Figure 2 are \( a = 6.380 \text{ Å}, b = 5.525 \text{ Å}, c = 5.189 \text{ Å} \) and \( a = 6.506 \text{ Å}, b = 5.634 \text{ Å}, c = 5.210 \text{ Å} \) respectively). Considering that the crystal structure and electronic structure (band component) are both similar to GaN (a natural consequence of the cation-mutation), we expect that various heterostructures (superlattice, core–shell quantum dots, etc.) based on LiAlGe₂N₄, LiGaGe₂N₄ and GaN (or even ZnO) can be readily fabricated, and flexible and continuous band structure engineering can be achieved in these lattice matched heterostructures, which may give rise to various applications in UV optoelectronic devices. Experimental synthesis of LiAlGe₂N₄ and LiGaGe₂N₄ is called for.

IV. CONCLUSIONS

Through cation-mutation in GaN, we found for the first time that two quaternary nitride semiconductors (LiAlGe₂N₄ and LiGaGe₂N₄) can be thermodynamically stable, whereas other quaternary nitrides I–III–V–I (I = Cu, Ag, Li, Na; K; III = Al, Ga, In) are not stable with respect to the phase separation into binary or ternary nitrides. Our results explain why quaternary nitrides had never been synthesized experimentally and indicate that the control of the Ga (Al) richness in the environment is critical for synthesizing single-phase LiGaGe₂N₄ (LiAlGe₂N₄). The calculated band structure and optical absorption spectra showed that the new quaternary nitrides (LiAlGe₂N₄ and LiGaGe₂N₄) are direct-band gap semiconductors with strong absorption of ultraviolet light. Furthermore, they are both lattice-matched to GaN or ZnO, and their valence band edge levels can be lower than that of ZnO; however, their conduction band edge levels are higher than that of GaN, and therefore, flexible band structure engineering can be achieved in the lattice-matched GaN/LiAlGe₂N₄/LiGaGe₂N₄ superlattices or heterostructures, which may work as high-performance ultraviolet optoelectronic devices.

■ ASSOCIATED CONTENT

1 Supporting Information The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.chemmater.5b03536.

Space group and formation energies of all considered secondary phases (PDF)

■ AUTHOR INFORMATION

Corresponding Author
*E-mail for S.C.: chensy@ee.ecnu.edu.cn.

Notes
The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

We thank Prof. Jiang Tang for the helpful discussion about the viable synthesis strategy. The work was supported by the National Natural Science Foundation of China (grant no. 91233121, 61125403), Shanghai Rising-Star Program (grant no. 14QJA1401500), Special Funds for Major State Basic Research (grant no. 2012CB921401, 2014CB921104), PCSIRT and the computer center of ECNU. P.N. is supported by the National Science Foundation and the Resnick Sustainability Institute at Caltech.

■ REFERENCES


