In situ microbeam surface X-ray scattering reveals alternating step kinetics during crystal growth

Guangxu Ju1,6✉, Dongwei Xu1,2, Carol Thompson3, Matthew J. Highland4, Jeffrey A. Eastman1, Weronika Walkosz5, Peter Zapol1 & G. Brian Stephenson1

The stacking sequence of hexagonal close-packed and related crystals typically results in steps on vicinal {0001} surfaces that have alternating A and B structures with different growth kinetics. However, because it is difficult to experimentally identify which step has the A or B structure, it has not been possible to determine which has faster adatom attachment kinetics. Here we show that in situ microbeam surface X-ray scattering can determine whether A or B steps have faster kinetics under specific growth conditions. We demonstrate this for organo-metallic vapor phase epitaxy of (0001) GaN. X-ray measurements performed during growth find that the average width of terraces above A steps increases with growth rate, indicating that attachment rate constants are higher for A steps, in contrast to most predictions. Our results have direct implications for understanding the atomic-scale mechanisms of GaN growth and can be applied to a wide variety of related crystals.

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1 Materials Science Division, Argonne National Laboratory, Lemont, IL, USA. 2 School of Energy and Power Engineering, Huazhong University of Science and Technology, Wuhan, Hubei, China. 3 Department of Physics, Northern Illinois University, DeKalb, IL, USA. 4 X-ray Science Division, Argonne National Laboratory, Lemont, IL, USA. 5 Department of Physics, Lake Forest College, Lake Forest, IL, USA. 6 Present address: Lumileds Lighting Co., San Jose, CA, USA.

✉ Email: juguangxu@gmail.com; stephenson@anl.gov
Our understanding of crystal growth is built on a powerful paradigm quantified by Burton, Cabrera, and Frank (BCF)\textsuperscript{1,2}, in which atoms are added to the growing crystal surface by attachment at the steps forming the edges of each exposed atomic layer, or terrace. The BCF model was originally developed for crystals with step heights of a full unit-cell and step properties that are identical from step to step for a given step direction. However, from the beginning\textsuperscript{3} it was recognized that there could be more complex situations. When the space group of the crystal includes screw axes or glide planes, the growth behavior can be fundamentally different on facets perpendicular to one of these symmetry elements\textsuperscript{2}. In this case, the terraces can still all have the same atomic arrangement, but now have different in-plane orientations of their top layer. The fractional-unit-cell-height steps that separate these terraces have structures and properties that can vary from step to step, even for a fixed step direction. Thus, surface morphologies with alternating terrace widths can arise that depend upon the deposition or evaporation conditions, as indicated in Fig. 1. The inequivalent kinetics at steps affects not only surface morphology but also the incorporation of alloying elements during crystal growth\textsuperscript{4,5}.

A ubiquitous but subtle version of this effect occurs on the basal-plane \{0001\}-type surfaces of crystals having hexagonal close-packed (HCP) or related structures, which are normal to a \(6\_\perp\) screw axis. Such crystals are made up of closely packed layers with 3-fold symmetry that alternate between opposite orientations, as shown by the \(\alpha\) and \(\beta\) terrace structures in Fig. 2c. On a vicinal surface, the \(\alpha\Theta\beta\) stacking sequence typically results in half-unit-cell-height steps. The lowest energy steps are normal to \{01\(T\)0\}\(-\)type directions, and have alternating structures conventionally labeled \(A\) and \(B\)\textsuperscript{6,9} as shown in Fig. 2a, b. When the in-plane azimuth of a step changes by 60°, for example, from \([01\bar{T}0]\) to \([10\bar{T}0]\), its structure changes from \(A\) to \(B\) or \(B\) to \(A\).

The alternating nature of the steps on such surfaces has been imaged in several systems, including Si\textsubscript{C}\textsuperscript{4}, GaN\textsuperscript{8,10–13}, AlN\textsuperscript{14}, and ZnO\textsuperscript{15}. These systems typically show a tendency for local pairing of steps (i.e., alternating step spacings), and an interlaced structure in which the step pairs switch partners at corners where their azimuth changes by 60°, as shown in Fig. 2d. These features are consistent with predictions that \(A\) and \(B\) steps have significantly different attachment kinetics\textsuperscript{6,8,12,16–22} that lead to unequal local fractions of \(\alpha\) and \(\beta\) terraces during growth. However, it has not been possible to experimentally distinguish the terrace orientation or step structure, and thus to determine whether \(A\) or \(B\) steps have faster kinetics.

In particular, the properties of \(A\) and \(B\) steps on GaN (0001) surfaces have been a matter of some disagreement. A seminal study\textsuperscript{8} of molecular beam epitaxy (MBE) of GaN observed alternating step shapes and proposed that the kinetic coefficients for adatom attachment are higher for \(A\) steps than \(B\) steps, that is, \(A\) steps grow faster for a given supersaturation. The support for this highly cited prediction is based on an argument regarding the difference in dangling bonds between \(A\) and \(B\) steps, and a comparison with experimental results on GaAs (111)\textsuperscript{23,24}. (Such face-centered cubic materials have \(A\)- and \(B\)-type steps that do not alternate between successive terraces and thus can be distinguished by their orientation\textsuperscript{9}.) In contrast, several subsequent theoretical studies of GaN (0001) organo-metallic vapor phase epitaxy (OMVPE) and MBE have consistently predicted that \(A\) steps have smaller adatom attachment coefficients than \(B\) steps. Kinetic Monte Carlo (KMC) studies of GaN (0001) growth under OMVPE conditions found step pairing\textsuperscript{17} driven by faster kinetics at \(B\) steps than \(A\) steps\textsuperscript{16}. The standard bond-counting energetics used in a KMC study of growth on an HCP lattice\textsuperscript{19} result in a much lower Ehrlich–Schwoebel (ES) barrier at \(B\) steps than at \(A\) steps, when only nearest-neighbor jumps are allowed. A recent KMC study of GaN (0001) growth under MBE conditions\textsuperscript{18} found triangular islands that close analysis reveals are bounded by \(A\) steps, indicating faster growth of \(B\) steps. An analysis of InGaN (0001) growth by MBE\textsuperscript{6} concluded that adatom attachment at \(B\) steps is faster, converting them into crenelated edges terminated by \(B\) steps.

The difference between the kinetics at \(A\) and \(B\) steps is a reflection of the chemical states of the adatoms, steps, and terraces that affect the dynamics of \(A\) and \(B\) steps. Studies of islands on the FCC Pt (111) surface\textsuperscript{25,26} have found that \(A\) steps have a higher growth rate than \(B\) steps, but that this relationship is reversed by the presence of adsorbates such as CO. An experimental study of AlN (0001) surfaces grown by OMVPE\textsuperscript{14} found a change in the terrace fraction as a function of the V/III ratio used during growth. Ab initio calculations of kinetic barriers on GaN and AlN (0001) under MBE and OMVPE conditions\textsuperscript{20–22} found that the barriers and adsorption energies at \(A\) and \(B\) steps depend

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**Fig. 1 Schematic of microbeam surface X-ray scattering during organo-metallic vapor phase epitaxy (OMVPE) growth.** Crystal truncation rod (CTR) measurements are sensitive to the \(\alpha\) or \(\beta\) terrace fraction changes that occur on vicinal (0001) surfaces of HCP-type crystals during deposition or evaporation. Calculated reflectivities are shown for the CTRs from the \((0\bar{1}1\bar{1})\) and \((0\bar{1}12)\) Bragg peaks (red and blue curves, respectively) with \(\alpha\) terrace fractions \(f_\alpha = 0.1\) and 0.9 typical for evaporation and deposition of GaN by OMVPE, as will be shown below.
in detail on the surface reconstruction induced by the environment. Thus, to properly understand, model, and control (0001) surface morphology in HCP-type systems, there is a need for an in situ experimental method that can distinguish adatom attachment kinetics at A and B steps in the relevant growth environment.

Here, we show that in situ surface X-ray scattering can distinguish the fraction of the surface covered by α or β terraces during growth, unambiguously determining differences in the attachment kinetics at A and B steps. X-rays are an ideal probe since they are sensitive to atomic-scale structure and can penetrate the growth environment. This method is enabled by using a micron-scale X-ray beam that illuminates a surface region of a high-quality single crystal having a uniform step azimuth, as shown in Fig. 1. We demonstrate this for OMVPE of (0001) GaN, with measurements of crystal truncation rods (CTRs) carried out in situ during growth. CTRs are streaks of intensity extending in reciprocal space away from every Bragg peak in the direction normal to the crystal surface, which are sensitive to the surface structure. We fit calculated CTRs from a model structure to these measurements to obtain the variation of the steady state α terrace fraction \( f_\alpha \) as a function of growth conditions, as well as the relaxation times \( t_{\text{rel}} \) of \( f_\alpha \) upon changing conditions. These results are compared to calculated dynamics based on a BCF model for a system with alternating step types to quantify the differences in the attachment rates at A and B steps.

Results

Calculated surface X-ray scattering with alternating step types. We present calculations showing how the intensity distribution along the CTRs is sensitive to the fraction of the surface covered by α or β terraces. Our calculations include the effect of surface reconstruction, using relaxed atomic coordinates that have been obtained previously. For a vicinal surface, the CTRs are tilted away from the crystal axes, so that the CTRs from different Bragg
peaks do not overlap. The X-ray reflectivity along the CTRs can be calculated by adding the complex amplitudes from the substrate crystal and the reconstructed overlayers, with proper phase relationships. Details of our calculations are given in a separate paper. Figure 3a shows calculated intensity distributions along (01T1) and (10T1) CTRs for the GaN (0001) surface, demonstrating how their L-dependences vary with \( f_\alpha \). Bragg peak locations have integer indices \( H_x K_y L_z \) in reciprocal lattice units; these indices identify the CTR associated with each peak. For this comparison, we use the 3H(T1) surface reconstruction and a fixed surface roughness, independent of \( f_\alpha \), as discussed in the “Methods” section below and Supplementary Discussion 1. The same qualitative behavior is obtained using other surface reconstructions. For \( f_\alpha = 0 \) and \( f_\alpha = 1 \), the regions between the Bragg peaks have alternating stronger and weaker intensities, with the alternation being opposite for (01T1) and (10T1). For \( f_\alpha = 0.5 \), the intensities between all Bragg peaks are about the same, and there is no difference between the (01T1) and (10T1) CTRs. As required by symmetry, the (01T1) CTRs with \( f_\alpha = X \) are identical to the (10T1) CTRs with \( f_\alpha = 1 - X \), for any value X. Figure 3b shows calculations of the reflectivity as a function of \( f_\alpha \) at positions near \( L = 1.6 \) on the (01T2) and (10T2) CTRs. The variation in reflectivity is almost monotonic in \( f_\alpha \) at these positions. These curves are used below to extract \( f_\alpha(t) \) during dynamic transitions.

In situ X-ray scattering measurements during growth. We studied four OMVPE conditions having different net growth rates at the same temperature 1076 ± 5 K, summarized in Table 1 (see “Methods” for further details). The substrate used was a GaN single crystal. Figure 4a shows its initial surface morphology determined by ex situ atomic force microscopy (AFM). One can see straight steps almost perpendicular to the \( y \) or (01T0) direction over large areas. An analysis of the step spacing shows a slight tendency towards pairing, with one of the two alternating terrace types having an area fraction of 0.47. AFM is insensitive to whether this fraction corresponds to the \( \alpha \) or \( \beta \) terraces. We also characterized the miscut angle by measuring the splitting of the CTRs. Figure 4b shows a transverse cut through the CTRs in the \( Q_y \) direction near (000L) at \( L = 0.9 \). Both the AFM and X-ray measurements give a double-step spacing of \( w = 573 \) Å corresponding to a miscut angle of 0.52°. To relate the \( \alpha \) terrace fraction to the behavior of \( A \) and \( B \) steps, it is critical to determine the sign of the step azimuth. By making measurements as a function of \( L \), we verified that the peak at high \( Q_y \) is the CTR coming from (0000), while the peak at low \( Q_y \) is the (0002) CTR. This confirms that the downstairs direction of the vicinal surface is in the \( +y \) direction, as drawn in Fig. 1. It is also useful to know the precise angle of the step azimuth with respect to the crystal planes, which determines the minimum kink density and thus the predicted values of some kinetic coefficients. X-ray measurements found this to be 5° off of the (01T0) direction towards (10T0), which gives a maximum kink spacing of 33 Å. The kink spacing could be smaller due to thermally generated kinks, as discussed in Supplementary Discussion 2. With this low-dislocation-density substrate and the low growth rates used, the previously reported instability to step bunching during growth was not observed.

Figure 5 shows the measured steady state CTR intensities as a function of \( L \), for both the (01T1) and (10T1) CTRs and at all four conditions. The qualitative behavior agrees with that
expected from a variation in \( f_a \) shown in Fig. 3a, with alternating higher and lower intensities between the Bragg peaks under some conditions, and opposite behavior of the two CTRs.

**Steady state and dynamics of terrace fraction during growth.**

To obtain values of the steady state terrace fraction \( f_a^{ss} \) for each of the four conditions, we fit calculated CTR intensities to the measured profiles. We performed fits using different possible surface reconstructions (see “Methods” for further details). While the 3H(T1) reconstruction gives the best fit to all conditions, similar \( f_a^{ss} \) values are obtained using alternative reconstructions. For each condition, both the (01T) and (10T) CTRs were simultaneously fit. The values of \( f_a^{ss} \) obtained as a function of net growth rate \( G \) are given in Table 1. The marked increase in \( f_a^{ss} \) as \( G \) is increased reveals the qualitative difference between the kinetics at \( A \) and \( B \) steps during OMVPE of GaN: adatom attachment coefficients for \( A \) steps are larger. Thus, a surface with initially balanced \( A \) and \( B \) terrace fractions at zero net growth rate will evolve to one with higher \( f_a^{ss} \) during positive net growth, because of the initially higher adatom attachment rate at the \( A \) steps. Likewise, during evaporation the initially higher detachment rate at \( A \) steps will give a lower \( f_a^{ss} \).

We also observed the dynamics of the change in \( f_a \) by recording the intensity at a fixed detector position as a function of time before and after an abrupt change between conditions, as shown in Fig. 6a. We chose positions near \( L = 1.6 \) where the X-ray reflectivity \( R \) changes almost monotonically with \( f_a \), as shown in Fig. 3b. It is thus straightforward to obtain \( f_a(t) \) from the intensity evolution using the calculated \( R(f_a) \), as shown in Fig. 6b. The characteristic 1/\( e \) relaxation times \( t_{rel} \) were 2200 \( \pm \) 200 and 340 \( \pm \) 30 s for the transitions from conditions 1 to 2 and 2 to 4, respectively.

**BCF model for surface with alternating step types.**

To quantitatively relate the behavior of the terrace fraction to the kinetic properties of \( A \) and \( B \) steps, we have developed a model based on BCF theory. Such models have been used extensively to understand growth behavior such as the step-bunching instability, pairing of steps, and competitive adsorption, typically where all steps in a sequence are equivalent. In our model, we consider an alternating sequence of two types of terraces, \( A \) and \( B \), and two types of steps, \( A \) and \( B \), with properties that can differ, as shown in Fig. 7. Related BCF models with an alternating step or terrace properties have appeared previously. We include the effects of step transparency (i.e., adatom transmission across steps) and step–step repulsion.

The rate of change in the adatom density per unit area \( \rho_i \) on terrace type \( i = A \) or \( B \) is written as

\[
\frac{\partial \rho_i}{\partial t} = D V_j^2 \rho_i - \rho_i + F, \tag{1}
\]

where \( D \) is the adatom diffusivity, \( r \) is the adatom lifetime before evaporation, and \( F \) is the deposition flux of adatoms per unit time and area. The four boundary conditions for the flux at the steps terminating each type of terrace can be written as

\[
-DV^A_i \rho^+_j + \kappa^A_j (\rho^+_j - \rho^A_j) + \kappa^A_0 (\rho^+_0 - \rho^B_0); \tag{2}
\]

\[
-DV^B_i \rho^-_j - \kappa^B_j (\rho^-_j - \rho^B_j) - \kappa^B_0 (\rho^-_0 - \rho^A_0); \tag{3}
\]

\[
-DV^B_i \rho^-_j + \kappa^B_j (\rho^-_j - \rho^B_j) + \kappa^B_0 (\rho^-_0 - \rho^B_0); \tag{4}
\]

\[
-DV^A_i \rho^+_j - \kappa^A_j (\rho^+_j - \rho^A_j) - \kappa^A_0 (\rho^+_0 - \rho^A_0). \tag{5}
\]

As shown in Fig. 7, \( \kappa^A \) and \( \kappa^B \) are the kinetic coefficients for adatom attachment at a step of type \( j = A \) or \( B \) from below or above, respectively, and \( \kappa^A_0 \) is the kinetic coefficient for transmission across the step. The + or − superscripts on \( \rho_i \) and \( V \rho_i \) indicate evaluation at the downhill or uphill terrace boundaries, respectively. We consider the overall vicinal angle of the surface to fix the sum \( w \) of the widths of \( A \) and \( B \) terraces, which are thus \( f_a w \) and \( (1-f_a) w \). We also assume relations between the equilibrium adatom densities \( \rho_i^{eq} \) at the steps and the terrace widths that reflect an effective repulsion between the steps owing to entropic and strain effects,

\[
\rho_i^{eq} = \rho_i^{eq} \exp(\mu_j/kT), \tag{6}
\]

where \( \rho_i^{eq} \) is the equilibrium adatom density at zero growth rate, and the adatom chemical potentials \( \mu_j \) at steps of type \( j = A \) or \( B \) have a dependence on \( f_a \) given by

\[
\frac{\mu_A}{kT} = -\frac{\mu_B}{kT} = M(f_a) \equiv \left( \frac{\ell}{w} \right)^3 \left[ \left( 1 - f_a^2 \right)^3 - \left( \frac{f_a}{f_a^2} \right)^3 \right], \tag{7}
\]

where \( \ell \) is the step repulsion length and \( f_a^2 \) is the terrace fraction at zero growth rate.
To solve this model, we develop a quasi-steady state expression for the dynamics of the terrace fraction $f_\alpha$. Under fairly general assumptions, the behavior of $f_\alpha$ can be written as a function of the net growth rate,

$$G = \frac{F - \rho_0^{eq}/\tau}{\rho_0}. \tag{8}$$

This is simply the difference between the deposition $F$ and a uniform evaporation $\rho_0^{eq}/\tau$, converted to monolayer per second (ML s$^{-1}$) using the site density $\rho_0$ per half-unit-cell-thick ML.

The rate of change in $f_\alpha$ is

$$\frac{df_\alpha}{dt} = K^{\text{dyn}}(f_\alpha) \left( \frac{G}{K^{\text{ss}}(f_\alpha)} - \frac{4M(f_\alpha)\rho_0^{eq}}{\nu p_0} \right), \tag{9}$$

where we have introduced the net steady state and dynamic kinetic coefficient functions $K^{\text{ss}}(f_\alpha)$ and $K^{\text{dyn}}(f_\alpha)$, which in the general case depend on all six $\kappa_j$ coefficients.

**Fig. 5 Measured and calculated crystal truncation rods.** Symbols show measured net intensities of the (01T\(_0\)) CTRs and the (10T\(_0\)) CTRs families (left and right) for CTRs from the $L_0 = 0, 1, 2$, and 3 Bragg peaks at each of four conditions. Curves show fits of all CTRs to obtain steady state $\alpha$ terrace fraction $f_\alpha^{ss}$ at each condition.
Fig. 6 Dynamics following a change of condition at t = 0. a Measured CTR intensities. b Calculated terrace fractions $f_a$. Blue curves: conditions 1-2 on the (10\(\overline{1}2\)) CTR at $t = 1.627$. Red curves: conditions 2-4 on the (01\(\overline{1}2\)) CTR at $t = 1.603$. Circles show 1/e relaxation times $\tau_{rel}$ of 2200 ± 200 and 340 ± 30 s, respectively.

Fig. 7 Schematic of alternating terraces and steps for the BCF model.

Vicinal (0001) surfaces of HCP crystals have alternating $\alpha$ and $\beta$ terraces separated by $A$ and $B$ steps. Notations are indicated for the kinetic coefficients for adatom attachment from below and above and for adatom transmission. Schematic of alternating terraces and steps of BCF model for HCP basal-plane surfaces, showing kinetic coefficients for the $A$ and $B$ steps.

The full steady state $df_a/dt = 0$ is obtained at a growth rate of

$$G^{\alpha}(f_a) = \frac{4K^{\alpha}(f_a) M(f_a)}{w\rho_0}.$$  

This equation for $G^{\alpha}(f_a)$ can be inverted to obtain the steady state value $f^{\alpha}_a$ as a function of $G$. The sign of $dG^{\alpha}/df_a$ and thus $df^{\alpha}_a/dG$ is determined by the sign of $K^{\alpha}$. General expressions for $K^{\alpha}$ show that in most cases, such as that found here, its sign is positive when the kinetic coefficients for $A$ steps are larger than those for $B$ steps.

To calculate BCF model results to compare with the experimental conditions, we make three assumptions: (1) the only parameter affected by the TEGa supply rate is the deposition flux $F$; (2) the only parameter affected by the carrier gas composition (0 or 50% H$_2$) is the adatom lifetime $\tau$; and (3) $F$ and $\tau$ enter only through the net growth rate $G$ given by Eq. (8), listed in Table 1 for each condition. We use the known values $\rho_0 = 2a^2/\sqrt{3} = 1.13 \times 10^{19}$ m$^{-2}$ and $w = c/\sin(0.52^\circ) = 5.73 \times 10^{-8}$ m, where $a = 3.20 \times 10^{-10}$ m

$$c = 5.20 \times 10^{-10}$$ m are the lattice parameters of GaN at the growth temperature. We performed fits to the measured quantities (four $f^{\alpha}_a$ and two $\tau_{rel}$) using the general expressions for $K^{\alpha}(f_a)$ and $K^{\alpha}(f_a)$. The relaxation times for the model were calculated by integrating Eq. (9) to obtain $f_a(t)$, and then extracting the relaxation time with the same normalization procedure used for the experimental data. The best fit, shown in Fig. 8, was obtained in the limit in which the $\kappa^\alpha$, $\kappa^\beta$, and $\kappa^\beta$ coefficients approach zero. In this case, $K^{\alpha}(f_a)$ and $K^{\alpha}(f_a)$ can be written as

$$K^{\alpha}(f_a) = \frac{1}{K_0^\alpha} + \frac{(1 - 2f_a)}{\kappa_0^\beta} - \frac{w(f_a)}{D}$$

and

$$K^{\beta}(f_a) = \frac{1}{K_0^\beta} + \frac{w(f_a)}{D}.$$  

Fig. 8 Steady state $\alpha$ terrace fraction $f^{\alpha}_a$ as a function of growth rate $G$. Circles are experimental values given in Table 1, showing monotonic increase of $f^{\alpha}_a$ with increasing $G$. Curve is best-fit BCF model calculation; also fit simultaneously with these four points were the two relaxation times $\tau_{rel}$ given in Table 1.

Note that $\kappa^\beta_0$ does not appear in these expressions because in this limit other microscopic processes are rate-limiting. We can fit the measurements directly using these expressions to obtain the four parameters $D/\kappa_0^\beta = 1.9 \times 10^{-8}$ m, $D/\kappa_0^\alpha = 1.1 \times 10^{-3}$ m, $D\rho_0^\alpha = 3.3 \times 10^{-23}$ m$^3$ s$^{-1}$, and $f^{\beta}_0 = 0.44$. Table 1 compares the six measured quantities (four $f^{\alpha}_a$ and two $\tau_{rel}$) to the best-fit values calculated from the BCF model.

Discussion

To interpret the combined parameters obtained from the fits, it is useful to estimate the adatom diffusivity $D$ and equilibrium adatom density $\rho_0^\alpha$. Ab initio calculations of the activation energy for Ga diffusion on the Ga-terminated (0001) surface have given values of $\Delta H_m = 0.44$ eV and 0.5 eV, and similar values have been obtained for 3d transition metal adatoms. If we estimate the diffusivity from the ab initio calculations using

$$D = a^2\nu \exp(\Delta S_m/k) \exp(-\Delta H_m/kT),$$

where $a = 3.2 \times 10^{-10}$ m, $\nu = 10^{13}$ s$^{-1}$, $\Delta S_m = 0$, and $\Delta H_m = 0.4$ eV, we obtain $D = 1.4 \times 10^{-8}$ m$^2$ s$^{-1}$ at $T = 1073$ K. An analysis of spatial correlations in surface morphology of GaN films indicated a cross-over at $T = 1073$ K from surface diffusion transport to evaporation/condensation transport at a length scale of $\lambda = 1.5 \times 10^{-6}$ m for OMVPE growth with H$_2$ present in the carrier gas. Thus, the adatom
lifetime $\tau$ can be estimated as $\tau = \lambda^2 / D = 1.7 \times 10^{-4} \text{s}$ under these conditions. Using our observed negative net growth rate for $F = 0$ of $G = -\rho_{eq}^0 (\rho_{0} \tau) = -0.00184 \text{ ML s}^{-1}$, this gives a value for the equilibrium adatom density of $\rho_{eq}^0 = 3.4 \times 10^{12} \text{ m}^{-2}$. Using these estimates for $D$ and $\rho_{eq}^0$, the parameters obtained from the mixed kinetics fit imply kinetic coefficients of $\kappa^a_0 = 0.74 \text{ m s}^{-1}$ and $\kappa^b_0 = 1.3 \text{ m s}^{-1}$, and a step repulsion length of $\ell = 9 \times 10^{-10} \text{m}$.

Although it has not been possible to use scanning-probe microscopy to observe the orientation difference of $\alpha$ and $\beta$ terraces on vicinal basal-plane surfaces of HCP-type systems, our results show that this difference is robustly revealed by surface X-ray scattering. In situ X-ray measurements during growth can determine the fraction covered by each terrace, and thus distinguish the dynamics of $A$ and $B$ steps. While the CTR calculations presented here are for wurtzite-structure GaN, this method applies to many other HCP-type systems with a 63 screw axis, minimum average kink spacing on the steps to be 3.3, and that the equilibrium adatom density of $\rho_{eq}$ is close to the saturation, and that the minimum average kink spacing on the steps to be 3.3. We expect that this will have important practical implications for advanced GaN devices, for example, laser diodes and white LEDs, by allowing control of interface morphology and incorporation of alloying elements such as In.$^2$.

We have demonstrated this method using a micron-scale X-ray beam to satisfy the requirement that the illuminated surface region has a well-defined step azimuth. With current synchrotron X-ray sources, it is possible to perform this type of experiment with highly monochromatic beams, greatly increasing the in-plane resolution of the CTR measurements.

### Methods

**Calculated CTR intensities.** Derivation of the CTR calculations for miscut surfaces with alternating terrace terminations is provided in a separate paper.$^{39}$ As described in Supplementary Discussion 1, in fits shown in Table 2, and in recent predictions$^{47}$, we expect that the GaN surface under OMVPE conditions has a 3H(T1) reconstruction, in which 3 of every 4 Ga atoms in top-layer sites shown in Fig. 3 are bonded to an adsorbed hydrogen. The calculations in Fig. 3a include the effect of this reconstruction, with equal fractions of all reconstruction domains on both terraces. Since no fractional-order diffraction peaks from long-range-ordered reconstructions are observed in experiments, we expect that the domain structure.

**Table 2 Values of fit parameters for each of four OMVPE conditions.**

| Growth condition index | Reconstruction | $f_a$ | $\sigma_\chi$ | $\chi^2$ |
|------------------------|----------------|------|--------------|--------|
| 1                      | 3H(T1)         | 0.111 | 0.75         | 106    |
|                        | Ga(T4)         | 0.144 | 1.26         | 130    |
|                        | NH(H3) + H(T1) | 0.098 | 0.94         | 187    |
|                        | NH(H3) + NH$_2$(T1) | 0.106 | 0.88 | 200    |
|                        | NH(H3)         | 0.095 | 0.91         | 167    |
| 2                      | 3H(T1)         | 0.461 | 0.93         | 57     |
|                        | Ga(T4)         | 0.476 | 1.26         | 81     |
|                        | NH(H3) + H(T1) | 0.460 | 1.15         | 76     |
|                        | NH(H3) + NH$_2$(T1) | 0.460 | 1.11 | 67     |
|                        | NH(H3)         | 0.459 | 1.13         | 99     |
| 3                      | 3H(T1)         | 0.811 | 0.85         | 118    |
|                        | Ga(T4)         | 0.670 | 1.46         | 218    |
|                        | NH(H3) + H(T1) | 0.876 | 1.19         | 205    |
|                        | NH(H3) + NH$_2$(T1) | 0.869 | 1.15 | 248    |
|                        | NH(H3)         | 0.869 | 1.16         | 168    |
| 4                      | 3H(T1)         | 0.868 | 0.47         | 80     |
|                        | Ga(T4)         | 0.942 | 1.06         | 112    |
|                        | NH(H3) + H(T1) | 0.892 | 0.90         | 174    |
|                        | NH(H3) + NH$_2$(T1) | 0.879 | 0.85 | 220    |
|                        | NH(H3)         | 0.891 | 0.87         | 135    |

We list the values of steady-state terrace fraction $f_a$, surface roughness $\sigma_\chi$, and the goodness-of-fit parameter $\chi^2$ from fits to reactivity for each of five surface reconstructions.
has a short correlation length and all domains are present. We use a surface roughness of $\sigma_0 = 0.74$ Å to match the experimental fits described below.

**In situ microbeam X-ray experiments.** We performed in situ measurements of the CTRs in the OMVPE environment at the Advanced Photon Source beamline 12ID-D\(^{36}\). At an incidence angle of $2\theta$, the 10 μm X-ray beam illuminated an area of 10 x 300 μm. To obtain sufficient signal, we used a wide-bandwidth pink beam setup\(^{35,36}\). Details of the measurements and data analysis are given in Supplementary Methods 1. Two types of measurements were performed. We determined the steady state terrace fractions $f_{ss}$ under four different growth/evaporation conditions by scanning the detector along the (01T1) and (10T1) CTRs while continuously maintaining steady state growth or evaporation. We also observed the dynamics of $f_{ss}$ after an abrupt change in conditions.

Under the conditions studied, deposition is transport limited, with the deposition rate proportional to the supply of the Ga precursor (triethylgallium, TEGa) with a large excess of the N precursor (NH\(_3\)) constantly supplied. We investigated conditions of zero deposition (no supply of TEGa) as well as deposition at a TEGa supply of 0.033 μmol min\(^{-1}\). The NH\(_3\) flow in both cases was 2.7 s.l.p.m. or 0.12 mol min\(^{-1}\), and the total pressure was 267 mbar. The V/III ratio during deposition was thus $3.6 \times 10^6$. For both of these conditions, we studied two carrier gas compositions: 50% H\(_2\) and 0% H\(_2\) (1073 K). Based on an expected activation energy of ~0.4 eV for the transition from conditions 2 to 4). The resulting growth rates $G$ equals 1/μslpm, or 0.12 mol min\(^{-1}\). A final fit using Eqs. (11) and (12) for $K^0(f_{ss})$ and $K(f_{ss})$ involves only the four known quantities $D_1/\sigma_0^2$, $D_2/\sigma_0^2$, $\Delta\sigma$, and $f_{ss}^2$. A fit of these parameters to give the same solution as the nine parameter fit.

**Data availability.** Data supporting the findings of this study are available within the article and its Supplementary Information file and are available in electronic form from the corresponding author on reasonable request.

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### Author contributions

All authors contributed to initial discussions motivating the experiments and analysis. G. J., C.T., M.J.H., I.A.E., and G.B.S. developed the microbeam surface X-ray scattering method and carried out the measurements. W.W. provided calculated atomic coordinates for the various surface reconstructions. D.X., C.T., P.Z., and G.B.S. developed the CTR and BCF expressions used in the analysis. G.J. and G.B.S. analyzed the results. All coauthors contributed to drafting and editing the manuscript.

### Competing interests

The authors declare no competing interests.

### Additional information

**Supplementary information**

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