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Modelling fine-sliced three dimensional electron diffraction data with dynamical Bloch-wave simulations

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Abstract

Recent interest in structure solution and refinement using electron diffraction (ED) has been fuelled by its inherent advantages when applied to crystals of sub-micron size, as well as a better sensitivity to light elements. Currently, data is often processed using software written for X-ray diffraction, using the kinematic theory of diffraction to generate model intensities – despite the inherent differences in diffraction processes in ED. Here, we use dynamical Bloch-wave simulations to model continuous rotation electron diffraction data, collected with a fine angular resolution (crystal orientations of $\sim 0.1^\circ$). This fine-sliced data allows us to reexamine the corrections applied to ED data. We propose a new method for optimising crystal orientation, and take into account the angular range of the incident beam and varying slew rate. We extract observed integrated intensities and perform accurate comparisons with simulations using rocking curves for a (110) lamella of silicon 185 nm in thickness. $R_1$ is reduced from 26% with the kinematic model to 6.8% using dynamical simulations.
1. Introduction

Electron diffraction (ED) is currently enjoying increased attention and activity due to its ability to work with crystallites that are far smaller than can be tackled by X-ray diffraction (XRD) (Gemmi et al., 2019; Grüne et al., 2018). Structural solution utilising ED has dramatically increased since the turn of the century due to advances in computer control and detector development (Gemmi & Lanza, 2019) and the new methodologies that have been developed for structure solution are generally known by the term three-dimensional electron diffraction (3D-ED) (Gemmi et al., 2019). Just as in XRD, these techniques measure the direction and integrated intensity of many Bragg-diffracted beams from a crystal, which are then processed to deduce a unit cell, given Miller indices $hkl$ and observed intensities $I_{hkl}^{(obs)}$. These data are then used to produce a crystal model using structure solution methods. There are many differences between ED and XRD, (Gemmi & Lanza, 2019) including very different wavelengths and damage mechanisms, but the principal one which affects diffracted intensities is the strength of interaction with electrons roughly 10,000 times more likely to be scattered than X-rays (Spence, 2017; Xu & Zou, 2019). It is this aspect which allows ED to outperform XRD at the nanoscale, but unable to tackle macroscopic crystals. The methods are therefore complementary, and together they make a powerful combination (Yun et al., 2015).

Even at the nanoscale, multiple scattering is usual for electrons – it is essential to capture the interaction of a fast electron with even a single gold atom (Howie, 2014) – whereas single scattering usually dominates for XRD. Since structural refinement relies on minimising the difference between $I_{hkl}^{(obs)}$ and calculated values $I_{hkl}^{(calc)}$, it is therefore unsurprising that a fit to a single (kinematic) scattering model is poor for a method where multiple (dynamical) scattering dominates. Currently, some analyses of ED data for structural solution and refinement still use the well-established and relatively
sophisticated XRD software, despite the vastly different scattering processes involved. As a result, structural solution statistics from 3D-ED often appear much worse than that of XRD, even though the structures obtained seem reliable. To develop the field of ED further, it is necessary to improve the quality of fit by taking into account the differences between electron diffraction and other methods. As first demonstrated by (Palatinus et al., 2015a), and now firmly established by (Klar et al., 2021), significant improvements can be obtained when dynamical electron scattering effects are considered.

The differences between electrons and X-rays when used for structure solution can be relatively subtle in data that only contain integrated intensities, where each reflection is measured over a time interval during which the diffracted beam passes completely through its Bragg condition. This type of data is found in continuous-rotation electron diffraction (cRED) experiments, where each data frame typically covers one or more degrees of crystal rotation. However, electron detector technology has seen a significant improvement in both quantum efficiency and speed in recent years (Faruqi & McMullan, 2018; Paterson et al., 2020; Grüne & Mugnaioli, 2021), allowing ever greater amounts of data to be obtained. These fast pixelated detectors can provide data that has fine resolution both temporally and in scattering angle. cRED experiments may now have data frames covering a small fraction of a degree (Fröjd et al., 2020). The additional information in such fine-sliced data allows the detail of electron scattering processes to be observed more clearly and provides an opportunity to model them more comprehensively. In this work, we explore data taken with a crystal orientation resolution of $\sim 0.1^\circ$, in combination with Bloch-wave electron diffraction simulations. Our aim is to elucidate the most important experimental and modelling parameters that will be necessary in future electron diffraction methods.

The main reason for the continued adherence to a scattering model that is known
to be inadequate for ED is the relative difficulty of calculation for dynamical scattering in comparison with the kinematic model. In both models, the starting point for calculation of the diffracted intensity for reflection \( \mathbf{g} = hkl \) is the structure factor \( F_{hkl} \)

\[
F_{hkl} = \sum_{j=1}^{N} f_j(\theta_B) T_j \exp{(2\pi i \mathbf{g} \cdot \mathbf{r}_j)}, \tag{1}
\]

where \( f_j(\theta_B) \) is the atomic scattering factor evaluated at the Bragg angle \( \theta_B \), \( T_j \) the thermal factor and \( \mathbf{r}_j \) fractional atomic coordinates of the \( j^{th} \) atom, the sum taken over all \( N \) atoms in the unit cell. In the kinematic model, it is commonly stated that the structure factor \( F_{hkl} \) is proportional to the amplitude of the diffracted beam, i.e. intensity is proportional to \( I_{hkl}^{(\text{kin})} = |F_{hkl}|^2 = F_{hkl} F_{hkl}^* \), where * indicates complex conjugate. With tabulated scattering factors Eq. (1) can be evaluated almost instantaneously on even the most basic computer. In comparison, modelling dynamical scattering for ED to obtain \( I_{hkl}^{(\text{dyn})} \) requires solving Schrödinger’s wave equation for an electron travelling through the crystal, usually done using either the Bloch-wave method or a multislice wave scattering/wave propagation calculation. The structure factor enters the Bloch-wave calculation as elements in the scattering matrix, which contains all excited \( \mathbf{g} \)-vectors and their differences, but is present only indirectly in a multislice calculation. These models, which require significant computing resources and may be cluster or GPU-based, are widely used in more traditional electron diffraction work such as convergent beam electron diffraction (CBED) (Tsuda & Tanaka, 1995; Spence, 1993; Zuo & Spence, 2013) and in the simulation of transmission electron microscopy (TEM) and scanning TEM (STEM) images.

Prior to the development of 3D-ED methods an additional difficulty in ED resulted from the collection of static diffraction patterns, meaning that with parallel illumination almost all reflections were measured away from the Bragg condition. In this case, even minuscule changes in crystal orientation can produce large changes in
diffracted intensity and while dynamical modelling was possible (Jansen et al., 1998) it was not straightforward. The introduction of beam precession (Vincent & Midgley, 1994) allowed measurement of intensities integrated around a circular path in reciprocal space, which reduced the sensitivity to precise crystal orientation. However, dynamical modelling of intensities in individual ED patterns with beam precession then had to take account of the different integration path for each g-vector (Sinkler & Marks, 2010; Dudka et al., 2007). In order to solve a crystal structure in three dimensions, multiple precession ED measurements were obtained at different crystal orientations (Kolb et al., 2007). This method has had great success but still requires corrections for integration of different g-vectors (Palatinus et al., 2015b). In the current cRED method, integration is performed by rotation of the crystal in a similar manner to 'X-ray precession measurements' (in which the crystal is referred to as 'precessed' which is referring to crystal rotation, not in the case with ED the incident beam). Extraction of integrated intensities in cRED data with, or without, beam precession is possible using the data reduction program Process Electron Tilt Series PETS (Palatinus, 2011; Palatinus et al., 2015a) and PETS2 (Palatinus et al., 2019) with structure solution and refinement with dynamical modelling using the programs JANA and Dyngo (Petříček et al., 2014; Palatinus et al., 2015b; Palatinus et al., 2019; Klar et al., 2021).

In any quantitative experiment, measurements and calculations must meet at some point and in XRD it is most convenient for that point to be the structure factor. The correspondence between model and experiment is usually captured using an R-factor (see section 3). Although it is often said that diffracted X-ray intensities from a crystal are proportional to $|F_{hkl}|^2$ (Eq.1), in reality things are not so simple and many other factors need to be accounted for, which depend on both the experiment being performed and the material. These factors are usually considered to be experimental
and can be either measured or determined during refinement, but are separate from the crystal structure itself. As noted by (Ladd & Palmer, 1994) it can be helpful to list these corrections in the form of an equation, Eq. 2. This makes a distinction between raw experimental measurements of diffracted intensity $I_{hkl}^{(expt)}$ and the ‘observed’ intensities $I_{hkl}^{(obs)}$ that are suitable for comparison with calculations $|F_{hkl}^{(calc)}|^2$.

$$|F_{hkl}^{(obs)}|^2 \propto I_{hkl}^{(obs)} = M^{-1} A^{-1} E^{-1} S^{-1} G^{-1} B^{-1} L^{-1} p^{-1} \{I_{hkl}^{(expt)}\},$$ (2)

with corrections for polarisation $p$, Lorentz factors $L$, background $B$, fluctuations in the incident beam intensity $C$, geometry $G$, scaling $S$, extinction $E$, absorption $A$, and mosaicity $M$. Each correction in Eq. 2 will be different for each reflection $hkl$ and should be considered a rather general operation when applied to a full diffraction data set. Furthermore if, like extinction in XRD, a correction is determined iteratively during refinement, it could be argued that they form part of a scattering model and therefore should be applied to $I_{hkl}^{(obs)}$ in Eq. 2 rather than $I_{hkl}^{(expt)}$. For simplicity here, where we are concerned with the effect of these factors rather than their point of application during data processing, we consider any modification of data necessary to improve structure solution and refinement to be applied to the raw experimental data, resulting in a set of ‘observed’ intensities that depend only on the crystal structure and nothing else.

In XRD each reflection $hkl$ has a single well-defined integrated $I_{hkl}^{(obs)}$, so that if it is sampled multiple times, or there are symmetrically equivalent reflections, they can be merged into a single measurement with improved fidelity. However, in dynamical diffraction a reflection no longer has a single well-defined intensity (Spence, 1993), as illustrated by Fig. 1. This shows a Bloch-wave simulation of a $\bar{1}4 \bar{2} \bar{2}$ reflection from the silicon cRED data set (section 3) which, although kinematically forbidden, has intensities up to 10% of the incident beam intensity where pathways for multiple allowed reflections exist. Although this is an extreme example, it is not uncommon
for weak reflections to be affected in this way in ED. Thus an average intensity, taken either by merging multiple measurements, symmetrically equivalent reflections, or through the use of a precessed incident beam (Vincent & Midgley, 1994), will in general converge to some ill-defined value. Conversely, comparison of experimental data with a dynamical model in which each measurement is simulated individually gives a better match without any data merging (Palatinus et al., 2015b).

Fig. 1. An illustration of the range of intensities that are obtained for the kinematically-forbidden $-14\ 2\ 2$ reflection in silicon with a thickness of 180nm in a dynamical simulation. The image below shows the corresponding LACBED pattern (see section 3)

Before embarking on a structural refinement in which $r_j$ and $T_j$ in Eq. 1 are determined for each atom in the unit cell by minimising the difference between observed and calculated intensities, the many corrections in Eq. 2 must be applied to optimise the experimental input $|F_{hkl}^{(obs)}|$. At first sight, it might therefore appear that Eq. 1 deals with scattering theory while Eq. 2 deals only with experimental parameters, but this
is not strictly correct – absorption and extinction, for example, are scattering effects that depend upon the sample. For dynamical electron diffraction, we must reconsider the validity of Eq. 2 to account for the differences in ED vs XRD experiments. For electrons, we can discard the polarisation correction $p$ since electron beams are unpolarised, but each of the other terms has an equivalent in ED. Thus, before presenting our results from a continuous rotation electron diffraction (cRED) measurement, we briefly discuss each in turn.

i) Lorentz corrections $L$. Lorentz corrections ensure that a given reflection has the same integrated intensity in XRD irrespective of the way the crystal is rotated, i.e. they account for the different time spent in the vicinity of each Bragg condition during data collection. As purely geometrical corrections, they apply equally to XRD and ED (Zhang et al., 2010b). If beam precession is employed in a set of ED patterns taken with a static crystal the circular path in reciprocal space taken by the direct beam must be taken into account (Gjønnes, 1997; Zhang et al., 2010a) but if data is collected with a continuously rotating crystal, beam precession adds no additional geometrical correction factor to integrated intensities (Palatinus et al., 2019). In our cRED experiment, we measure the integrated intensity of each reflection individually from rocking curves in both experiment and simulation and it is therefore possible to compare intensities without applying Lorentz corrections. However, to maintain equivalence with XRD refinement methods it is preferable to apply them, and we do so for both experiment and simulation.

ii) Geometry $G$. Corrections for other geometrical issues, such as variations in specimen height or rotation axis, are needed in XRD and also for ED (Palatinus et al., 2019). These issues primarily affect a diffracted beam’s position, rather than the intensity that is our main interest here. It is of course important to ensure that the crystal of interest does not wander out of the electron beam as it is rotated (Cichocka
et al., 2018; Plana-Ruiz et al., 2020). More importantly, since dynamical diffraction is exquisitely sensitive to geometry, crystal orientation must be known to high precision. At crystal orientations where diffraction is strong, a good calculation of intensities in a cRED measurement using dynamical scattering requires an angular precision and accuracy better than 40 arc seconds (~0.2 mrad). Previously, orientation refinement has been performed by optimising the fit between experimental and calculated intensities, either integrated (Palatinus et al., 2013) or pixel-wise for peaks detected on individual frames (Palatinus et al., 2019). Here, we show that fine-sliced data allows a quicker and more straightforward orientation refinement to be performed, using the sequence of reflections as they appear during crystal rotation. The fine sliced data approach also permits a measurement of varying slew rate (section 3). As a corollary of the requirement for high precision in crystal orientation, the angular range of the incident beam must also be taken into account; while parallel illumination is often assumed, in practice there is always some beam convergence or divergence that broadens the range of reciprocal space that is sampled.

iii) Background B. In ED, corrections for background need to be considered from two main sources, a) the detector; and b) non-Bragg scattered electrons (both elastic and inelastic). For (a), detector characteristics such as dark noise levels, quantum efficiency and linearity can be measured effectively; these corrections are necessary but straightforward. Unfortunately, (b) is rather more problematic. Amorphous material, e.g. the support film for the crystal, can create a non–linear background (Tivol, 2010), but the sample itself also produces non-Bragg thermal diffuse scattering (TDS, caused by displacement of atoms from their mean positions by thermal vibrations) and inelastic scattering. Importantly, these electrons can be diffracted again by the crystal, producing a background that is highly structured, with Kikuchi lines and dynamical scattering effects (Eggeman & Midgley, 2012). Calculating this background is not a
trivial exercise and generally requires a model of thermal vibrations (ideally, complete knowledge of the phonon spectrum) (Muller et al., 2001; Kolb et al., 2012) and a full quantum-mechanical description of inelastic processes (Forbes et al., 2011) respectively. In a complete model of electron scattering these effects would be taken into account in the calculation of diffracted intensities to be compared against experiment. Currently, while some current multislice simulation packages can do so (Allen et al., 2015) none have yet been implemented for 3D-ED experiments. Here, we use a simple Bloch-wave model that neglects this ‘background’ intensity of diffuse scattering by the crystal.

iv) Absorption $A$. This is another term which, strictly speaking, should be considered in scattering theory but in XRD its behaviour is simple enough for it to be corrected as an experimental variable. For the energies typical of ED (80-300 keV) and a thin specimen suitable for structure solution, true absorption of the electron beam does not happen to any appreciable extent. However, the attenuation of a Bragg reflection, as electrons are scattered into the diffuse background by TDS or inelastic interactions, is a very similar effect. In ED, TDS is enhanced significantly when the electrons are channeled along atom columns, particularly those with high atomic number (Hall & Hirsch, 1965). Thus, dark bands can be seen between low-index Bragg conditions in bright field LACBED patterns (‘anomalous absorption’, (Hirsch et al., 1965; Jordan et al., 1991) see examples below). This complicated behaviour means that in ED it is best dealt with in scattering theory, and should no longer regarded as an experimental parameter. Both Bloch-wave and multislice models can account for this effect in ED.

v) Extinction $E$. This is simply the X-ray term for dynamical diffraction effects. The underlying theory is very similar; the two-beam analysis by Darwin (Darwin, 1914a; Darwin, 1914b; Darwin, 1922) for X-rays has many resemblances to that of Howie and Whelan for electrons (Howie & Whelan, 1961; Hirsch et al., 1965), so much so that
they are sometimes referred to as the Darwin-Howie-Whelan model (James, 1990). The term *extinction* refers to the transfer of intensity from the direct beam to a diffracted beam $\mathbf{g}$, and back again, as a function of crystal thickness; the distance over which this occurs is known as an extinction distance $\xi_g$. In XRD this is usually dealt with during refinement and can be considered as a multiplicative correction factor that depends only upon $F_{hkl}$ and $\mathbf{g}$-vector magnitude (Becker & Coppens, 1974; Petříček et al., 2014; Bourhis et al., 2015). This simple approach can be used because X-ray extinction distances are usually much larger than the size of a crystallite. In ED, where extinction distances can be tens of nm and multiple beams are almost always excited, this is not the case. (In XRD, there is also ‘secondary’ extinction, which refers to the enhanced absorption of strong diffracted beams in large crystals (Ladd & Palmer, 1994), whose counterpart in ED is anomalous absorption, above). Again, this correction should not be applied to ED data, but taken into account in the calculation of diffracted intensities.

vi) *Scaling S*. This correction takes account of the varying proportion of the incident beam occupied by a crystal of irregular shape as it is rotated. This is certainly a correction that should be applied in principle in ED, although in practice extreme care must be taken not to confound it with simple loss of intensity in the direct beam due to a very strong diffracted beam, or the effect of ‘absorption’ due to TDS. ED has a potential advantage over XRD here, in that a crystal can be imaged directly, allowing scaling to be calculated from a series of images taken after a diffraction measurement (Plana-Ruiz et al., 2020). Without these images, frame-by-frame scaling of intensities is difficult to calculate (e.g. the current implementation of dynamical refinement refines the scale as part of the refinement process rather than data reduction process (Palatinus et al., 2019)). Here we propose a method using the direct beam intensity.
vii) Fluctuations in incident electron beam intensity C. cRED measurements are in general very rapid; diffracted electron beam intensities are sufficiently high that they can be sampled to good precision even in a small fraction of a second and the total time for data collection is often less than a minute. Variations in incident beam intensity on this timescale are negligible. Despite this, it is common to see significant changes in the direct 000 beam intensity in a cRED dataset (e.g. SI video 1). This happens because electron diffraction is strong and the crystal occupies much, or all, of the incident electron beam. In ED, it is quite possible for a diffracted beam with a large structure factor to have a higher intensity than the direct beam. Nevertheless, this effect is not due to a change in incident beam intensity and therefore this correction is not appropriate for a cRED measurement.

vii) Mosaicity M. Crystal imperfections, in the form of dislocations, low-angle grain boundaries and cracks, allows the Bragg condition to be satisfied over a wider range of angles than would be the case for a perfect crystal without strain. In XRD, the presence of these defects can be helpful in that they effectively break the crystal into a mosaic of small crystal blocks that reduces extinction effects significantly, (Ladd & Palmer, 1994) although it can also produce broadening of diffracted beams. For high energy electrons, with much smaller extinction distances, defects alter diffracted intensity very strongly, allowing them to be visualised directly in diffraction-contrast TEM (Hirsch et al., 1965; Williams & Carter, 2009). In ED they may have a significant effect on measured intensities that is too complicated to account for in any simple model and will vary from one crystal to another in an unknown way. Currently, their effect is neglected completely and this is probably the best approach until the more tractable effects of dynamical diffraction are fully accounted for.

To summarise, in reconsidering Eq. 2 for dynamical ED structure solution or refinement, we find that some factors that can be separated from the scattering model
and considered ‘experimental’ variables for XRD – i.e. absorption $A$, geometry $G$, and extinction $E$ – must instead be considered explicitly in the scattering model for ED. Other experimental factors – polarisation $p$, incident beam intensity variations $C$, and the complicated effects of mosaicity $M$ – can probably be neglected at the current level of simulation fidelity, being either relatively unimportant or too difficult to tackle with current methods. Finally, truly experimental corrections $L$, $B$ and $S$ that both XRD and ED hold in common must be taken into account, but the differences in Bragg angle and hardware means that they are rather different for ED. One of the biggest changes in emphasis is that the point of contact between experimental measurements and modelled intensities in a dynamical refinement no longer has the simple and elegant interpretation relating to the structure factor.

In what follows, we use the reliability factor $R_1$ (see S1.3) as a metric to compare model and experiment, using the square root of the observed (corrected) intensity $I_{hkl}^{\text{obs}}/2$. Our discussion above justifies only corrections to the raw data for Lorentz factors $L$, background $B$ and scaling $S$:

$$I_{hkl}^{\text{obs}} = S^{-1} B^{-1} L^{-1} \{I_{hkl}^{\text{expt}}\}.$$  (3)

We compare $I_{hkl}^{\text{obs}}$ with the square root of calculated ED integrated intensities $I_{hkl}^{\text{calc}}$ using a dynamical diffraction model in which extinction $X$ and absorption $A$ are implicit, resulting from the crystal itself. Dynamical integrated intensities $I_{hkl}^{\text{dyn}}$ are a non-linear function of crystal thickness $t$, geometry $G$ and beam profile $P$. It should be borne in mind that the relationship between these values and the structure factor $F_{hkl}$ of Eq. 1 is no longer straightforward. Since we apply Lorentz correction factors $L$ to the simulation we distinguish between the raw simulation $I_{hkl}^{\text{dyn}}$ and corrected intensities $I_{hkl}^{\text{calc}}$:
\[ I_{hkl}^{(calc)} = L^{-1} \left\{ I_{hkl}^{(dyn)} (X, t, G, P) \right\}, \quad (4) \]

We now explore this approach using Bloch-wave simulations to compare kinematic and dynamical model fits to experimental data for a simple, well-known material. We examine data from a single crystal of silicon, ion milled to a thin foil. This almost perfect crystal shows strong dynamical scattering and is used to evaluate the improvement in fit using dynamical modelling as well as the importance of correction factors applied to the raw data.

2. Experiment

2.1. Data collection

Experimental data was obtained using selected area electron diffraction (SAED) with parallel beam illumination on a JEOL 2100 LaB$_6$ transmission electron microscope operating at 200 kV. The sample was a defect-free single crystal milled using precision ion polishing (PIPS) Ar$^+$ ions at 6 kV to electron transparency with surface damage minimised by final polishing at 0.5 kV, producing a (110) lamella with extensive thin, parallel-sided regions. Diffraction patterns were produced using a strongly excited third condenser lens giving close to parallel illumination and a selected area aperture, and captured using a Gatan OneView camera recording continuously at 8× binning (4096 × 4096 → 512 × 512) at 75 frames/sec. (N.B. with 8x binning for data collection, the point spread function of the camera is negligible.) The crystal was much larger than the selected area aperture and was rotated at maximum slew rate about the α-tilt axis over 143°, giving 1389 frames, each with a nominal crystal rotation of 0.1035° (i.e. all data collected in 18.5 seconds). The angular step per frame was determined by finding the lowest score parameter in PETS from the cylindrical projection plots.
2.2. Data reduction and dynamical simulations

We used PETS (Palatinus, 2011) to find the crystal orientation and unit cell, index reflections and give their rocking curves. The procedure followed sections 4.1-4.6 of (Palatinus et al., 2019), which refers to the newer version PETS2 (Palatinus et al., 2019) but remains accurate for both versions.

Bloch wave simulations were performed using the code Felix (Beanland et al., 2019) running on a high performance computing cluster (typically 384 cores, completing a simulation in $\sim 30$ seconds) using a python script to extract data from the the dyn.cif.pets file generated by PETS and write input files for Felix, e.g. setting the incident beam orientation with respect to the crystal. Large-angle convergent beam electron diffraction (LACBED) patterns of size 400x400 pixels were simulated over an angular range corresponding to 40 frames ($4.14^\circ$, or 72.26 mrad). The $x$-axis of the image was taken to be along a direction perpendicular to the rotation axis. Successive simulations overlapped by 10 frames in an attempt to ensure that each reflection was fully captured in a single simulation, an approach referred to as Overlapping Virtual Frames (OVF) (Klar et al., 2021). Seventy simulations were required in total to cover the full angular range of the cRED data.

3. Results

The Si cRED data contained 962 reflections up to 4Å$^{-1}$, with intensity over 6000 or $I_{hkl}^{(expt)}/\sigma_{hkl} > 2$. Almost all reflections showed a single sharp peak as the crystal was rotated; only 27 had clear dynamical structure with multiple peaks in their rocking curve, obtained as an intensity given frame-by-frame as an output from PETS. Although these data were nominally background corrected ($B^{-1}$ in Eq. 3), close inspection showed the tails of the rocking curves had a small non-zero average. An additional small background correction was therefore performed by subtracting a lin-
ear fit interpolated beneath the peak of each rocking curve (supplementary Fig. S1).

The origin of the small non-zero intensity when no peak was present is unclear, although we note that complex background from thermal diffuse scattering and Kikuchi lines are readily apparent in the raw data as shown in Fig. 2 (see also the animated dataset in supplementary information).

Fig. 2. Seven consecutive frames from the Si cRED data set, showing Kikuchi lines as diffuse contrast passing through Bragg spots, marked by magenta and yellow arrows.

The Lorentz correction $L^{-1}$ was then applied for each reflection $g$ by taking the sum of counts in the background-subtracted rocking curve and multiplying by the change in deviation parameter per frame $\delta s_g$,

$$I'_{hkl} = \delta s_g \sum I^{(frames)}_{hkl}.$$  \hspace{1cm} (5)

Since the silicon lamella was close to parallel-sided and much larger than the selected area aperture, it filled the field of view completely throughout data collection and therefore no frame scaling correction $S^{-1}$ should be required. Nevertheless, it is useful to examine the total intensity falling on the detector and the intensity of the direct beam as the crystal rotated, both shown in Fig. S2. The total intensity (blue line) shows a drop of approximately 10% when the crystal is perpendicular to the beam, close to zero $\alpha$ tilt. Since the silicon is not thick enough to absorb 200 kV electrons to any appreciable extent and the short collection time precludes any significant change in incident beam intensity, the change in total intensity must be due to scattering of electrons to high angles, outside the detector area. This observation remains
unexplained. For the direct beam, each frame was cropped to a \( x = 17 \times y = 17 \) pixel image \( I(x, y, z) \) containing just the direct beam. An average beam profile was obtained by summing all frames and dividing by \( n = 1387 \), i.e.

\[
\bar{I}_{000}(x, y) = \frac{1}{n} \sum_{z=1}^{n} I(x, y, z)
\]

A normalised direct beam intensity \( I'_{000} \) was then produced by dividing each cropped frame by \( \bar{I}_{000} \):

\[
I'_{000}(x, y, z) = \frac{I_{000}(x, y, z)}{\bar{I}_{000}(x, y)}
\]

Re-slicing this \( 17 \times 17 \times 1387 \) \( xyz \) data volume to \( 1387 \times 17 \times 17 \) \( zyx \), and taking the average along \( x \), gives a \( 1387 \times 17 \) \( yz \) image that shows relative direct beam intensity as the crystal is rotated (supplementary Fig. S2, black line in Fig. 3). The trend in total intensity is also visible here (orange line). There is a range of approx. 10° at \( \alpha = 0° \) where the relative intensity is depressed by \( \sim 20\% \), but this is primarily due to channeling (see below); since this is calculated as part of the dynamical simulation it should not be compensated by scaling. Most strikingly, there are many sharp and significant minima (up to 50% of the relative intensity). Closer examination showed each of these dark lines to occur when a diffraction condition was satisfied; they simply indicate a transfer of intensity from the direct beam to a diffracted beam. As seen in more detail in supplementary Fig. S2, Bragg conditions that are satisfied in only a few frames are visible as dark vertical bands, while Bragg conditions that pass through the direct beam slowly are visible as inclined dark lines. In the absence of any explanation of the broad variations in intensity and knowing that the crystal completely filled the selected area aperture, no intensity scaling was applied.
Fig. 3. Blue line: the total intensity in the Si cRED ED pattern as a function of goniometer angle $\alpha$, normalised to an average of unity and offset by $+0.4$. Black line: the relative direct beam intensity for the data, normalised to an average of unity (red line). See also Fig. S2.

After applying corrections $B^{-1}$, $L^{-1}$ and while discounting $S$, we perform an overall scaling to directly compare simulated and experimental intensities. Intensities are calculated with an incident beam intensity of unity and we multiply corrected integrated observed intensities by the factor $K$:

$$K = \frac{\sum f_{hkl}^{(\text{calc})} I_{hkl}^{(\text{obs})}^{1/2}}{\sum f_{hkl}^{(\text{obs})} I_{hkl}^{(\text{obs})}^{1/2}}$$

(8)

giving a set of integrated intensities $KI_{hkl}^{(\text{obs})}$ that can be compared to simulation and yield an $R$-factor. We use the usual definitions, e.g. (Palatinus et al., 2013):

$$R_1 = \frac{\sum \left|KI_{hkl}^{(\text{obs})} - f_{hkl}^{(\text{calc})}\right|^{1/2}}{\sum \left|KI_{hkl}^{(\text{obs})}\right|^{1/2}}$$

(9)

$$R_2 = \frac{\left(\sum \left|KI_{hkl}^{(\text{obs})} - f_{hkl}^{(\text{calc})}\right|\right)^2}{\sum KI_{hkl}^{(\text{obs})}}$$

(10)

$$wR_2 = \left[\sum w_{hkl} \left(KI_{hkl}^{(\text{obs})} - f_{hkl}^{(\text{calc})}\right)^2\right]^{1/2} \sum w_{hkl} KI_{hkl}^{(\text{obs})}$$

(11)

where the weights $w_{hkl} = 1/I_{hkl}^{(\text{obs})} \sigma_{hkl}^2$ (Ladd & Palmer, 1994). We first consider kinematic intensities calculated using Eq. 1, taking the temperature factor to be
\[ T = \exp(-B \sin^2(\theta)/\lambda^2) \] where \( B_j \) is the Debye-Waller factor. We find a minimum \( R_1 = 26.4\% \) \( (R_2 = 59.4\%, wR_2 = 56.2\%) \) at \( B = 0 \), shown in Fig. 4 (X-ray refinements give \( B = 0.54\text{Å}^2 \) (Többens et al., 2001)), In this plot the bold black line indicates \( R_1 = 0 \) \( (F_{hkl}^{\text{obs}} = F_{hkl}^{\text{kin}}) \) while the orange line is a least-squares linear fit to the data. It is quite clear that the fit of the kinematic model to the data is poor, not least because there are a number of kinematically–forbidden reflections that have significant experimental intensities (highlighted in red), but also because many strong reflections \( (F_{hkl}^{\text{kin}} > 0.2) \) are in fact weaker than expected.

![Graph showing R1 calculation for cRED silicon data using the kinematic model (Eq. 1) with B = 0.2. Kinematically forbidden reflections are highlighted in red. R1 = 26.0%](image)

The poor \( R_1 \) seen in Fig. 4 is typical of many ED refinements using a kinematic model and we now turn to a dynamical one. We use an initial Debye-Waller factor \( B = 0.54 \) (Többens et al., 2001) since the value obtained from the kinematic model is unphysical. A Bloch-wave simulation of seventy Felix 000 LACBED patterns, stitched together to make a continuous strip, is shown in Figure 5. Frame numbers underneath correspond to experiment. In this image, a perfect plane-wave incident beam corre-
responds to a single point and the red line marks the nominal path traced by the direct beam through reciprocal space as the crystal is rotated. Each dark line in the simulation shows the location of a Bragg condition; when the direct beam lies on one of these lines a diffracted beam is produced, with an intensity that can be obtained from the corresponding point in the relevant dark-field LACBED pattern. The correspondence between this simulation and experiment can also be seen by converting the normalised experimental direct beam data volume to a 2D image, shown in grey-scale below the simulation. Some features common to both experiment and simulation are marked by arrows.
Fig. 5. Dynamical simulation for the path of the 000 beam through reciprocal space (red line) in the silicon cRED data set, corresponding to 1387 frames (goniometer rotation of 143° about \( \alpha \)). Frame numbers are indicated; note that one division = 5 frames = 0.5175° = 9 mrad. Each dark line corresponds to a Bragg condition for a diffracted beam, two of which (\( \bar{9}\bar{3}\bar{3} \), frames 30-50 and 311, frames 545-590) are labelled. Below the simulation, the experimental intensity of the 000 beam is shown. Some features that are clearly present in both simulation and experiment are indicated by arrows. The [110] zone axis lies a few degrees below the beam path around frame 670.

The simulated rocking curve for a reflection is given by the intensity along a line in its dark field LACBED pattern, as shown in Fig. 6 for a) a typical reflection with a single peak \( \bar{9}\bar{3}\bar{3} \), and b) one with obvious dynamical structure 311. Integrated intensities
$I_{hkl}^{(dyn)}$ can be obtained from these simulated rocking curves in the same way as they are taken from experimental ones, although here there is no diffuse scattering and therefore no background correction. The incident beam intensity is fixed at unity so there is no scaling correction and only Lorentz corrections need to be applied.

Fig. 6. Two examples of silicon cRED rocking curves. (a) and (b) are experimental data. Most experimental rocking curves have a simple peak like (a) 933, while less than 3% show dynamical structure like (b) 311. c) and d) show the corresponding dark field LACBED simulations (specimen thickness 185 nm). The nominal beam path is a red line, with frame numbers in yellow. Intensity profiles along the red line give the rocking curves (e) and (f). The difference in frame numbers (a) to (e) and (b) to (f) are caused by a varying slew rate (Fig. 9b). g) and h): Applying the angular spread of the incident beam (Fig. 10) as a convolution to the simulation gives simulated rocking curves that are a good match to experiment.
Kinematic intensities are independent of crystal thickness, but dynamic intensities can be very sensitive to it, particularly strong reflections with short extinction distances. It is therefore necessary to perform simulations for a range of thicknesses. (Supplementary sections S1.4 and S1.5). We may also expect the specimen thickness along the path of the electron beam to vary as the crystal is rotated, but this is ignored for the moment. The best $R_1$ is obtained for a thickness of 190nm, shown in Fig. 7. A very significant improvement over the kinematic model is apparent, with $R_1 = 12.6\%$ ($R_2 = 23.8\%, wR_2 = 39.8\%$). The wide spread of intensities is no longer present, but there is still considerable scatter about the expected $R_1 = 0$ line and the gradient of a linear fit is 0.74.

![Fig. 7. $R_1$ calculation for cRED silicon data with a Bloch wave model, using the nominal direct beam path (red line in Fig. 3) and a specimen thickness of 190 nm. The kinematically forbidden reflections identified in Fig. 4 now have non-zero values. The black line marks perfect correspondence ($R_1 = 0$) and the orange line a least squares linear fit. $R_1 = 12.6\%$](image)

Fig. 7. $R_1$ calculation for cRED silicon data with a Bloch wave model, using the nominal direct beam path (red line in Fig. 3) and a specimen thickness of 190 nm. The kinematically forbidden reflections identified in Fig. 4 now have non-zero values. The black line marks perfect correspondence ($R_1 = 0$) and the orange line a least squares linear fit. $R_1 = 12.6\%$

Dynamical simulations therefore clearly give a much better fit to experimental data, as may be expected. However, $R_1$ is still relatively high in Fig. 7 and further improve-
ments are possible by increasing the precision of the crystal orientation. The wide range of reciprocal space covered in the simulation allows geometry to be optimised, as described below.

3.0.1. Orientation optimisation G. For any given reflection in Fig. 5 its Bragg condition is satisfied, and a spot will appear in the SAED pattern, when the 000 beam sits on the corresponding dark line. The frame in which the maximum diffracted intensity appears is given by the crossing point of the Bragg condition and the red line. It is thus possible to obtain the sequence of reflections which appear in a cRED experiment, and the frame spacing between them, for any given direct beam path. Conversely, with knowledge of the frames in which diffracted maxima appear in an experiment we can find the corresponding path through reciprocal space.

Fig. 8. a) The intersections with Bragg conditions for eight reflections in the first Si Felix simulation. Frame numbers are given at the bottom of the image and the frame in which each peak intensity is found is marked by a vertical blue line. The intersection of the blue line with its Bragg condition is marked by a yellow dot (yellow lines correspond to an error of ±0.5 frames). The horizontal red line marks the nominal beam path (output from PETS) and if the crystal orientation was correct the yellow dots would all lie on this line. b) The best fit to a straight line is obtained by shifting the blue lines by -2.5 frames. c) $R_1$ calculation using an optimised beam path and a specimen thickness of 185 nm. $R_1 = 10.0\%$

The effect of a slightly incorrect crystal orientation is shown in Fig. 8a. In this simulated image each frame corresponds to a vertical stripe ten pixels wide and the
nominal direct beam path runs horizontally through the centre, marked by a red line. Experimentally, the $\bar{1}4 \ 2 \ 2$ and $11 \ 1 \ 3$ reflections were seen in frame 4, $8\bar{4} \bar{4}$ was seen in frame 9, etc, as marked by blue vertical lines. The image is a superposition of eight dark-field LACBED patterns; each bright line corresponds to a diffracted beam (and to a dark line in the direct beam LACBED pattern, not shown). The intersection of each blue line and its corresponding diffraction condition is marked by a yellow dot and must lie on the beam path; yellow lines indicate an error of 0.5 frames. Clearly, these points do not correspond to the expected horizontal line through reciprocal space, indicating that the nominal crystal orientation is slightly in error. An optimised crystal orientation can be found by shifting the group of blue lines while maintaining their relative frame spacing (e.g. $3\bar{3} 9$ has an experimental peak intensity one frame before $13 \bar{7}$ and three frames after $8\bar{4} \bar{4}$). Shifting the set of blue lines by 2.5 frames to the left brings all points close to a horizontal line (Fig. 8b). An optimised direct beam path was then calculated by fitting a smoothed curve to the best crystal orientation for each simulation (Fig. 9), both for changes about the rotation axis $\delta \alpha$ and perpendicular to it $\delta \beta$ by least-squares fitting a horizontal line to the optimised set of intersection points. Rocking curves extracted from the Bloch-wave simulations using an optimised beam path gives a significant improvement to $R_1 = 10.0\%$ ($R_2 = 19.9\%$, $wR_2 = 30.2\%$, Fig. 8c), mainly by reducing the scatter in reflections with lower intensities. This can be understood by referring back to Fig. 1, which shows how large variations in intensity can be found in weak beams when they coincide with stronger beams. Optimisation of the beam path is essential to capture these interactions correctly.

The resulting corrections are shown in Fig. 9. The actual path traced by the direct beam deviates vertically from the red line (i.e. about the $\beta$ tilt axis) in Fig. 9 by a maximum of 10 pixels (equivalent to 1 frame, or $0.1^\circ$). There is a much larger
correction needed along $\alpha$, up to 5 frames ($0.5^\circ$, Fig. 9b) that varies through the data series, caused by a varying slew rate during rotation of the specimen. This varying slew rate is apparent in Fig. 5, where the features in the normalised experimental direct beam intensity are not found directly beneath their corresponding points in the simulation above. A changing slew rate also has an impact on integrated intensities, since the crystal is rotating more slowly or quickly through a diffraction condition than expected. Applying a slew rate correction to the simulated intensities gives a further improvement to $R_1 = 9.4\%$. ($R_2 = 13.8\%, wR_2 = 28.7\%$, Fig.9c).

Fig. 9. Corrections to the nominal direct beam path (red line in Fig. 5), a) perpendicular to the line ($\beta$-tilt) and b) along the line ($\alpha$-tilt). Uncertainty in $\delta\alpha$ and $\delta\beta$ for each data point are $\sim \pm 0.5$ frames, i.e. $0.05^\circ$. c) $R_1$ calculation after correcting integrated intensities for varying slew rate gives $R_1 = 9.4\%$ at a specimen thickness of 195 nm.

3.0.2. Correction for beam profile $P$. The sensitivity of dynamical electron diffraction to thickness is apparent in the $R_1$ calculation for all integrated intensities (see S1.4). It is also very important for the fine structure of rocking curves of strongly dynamical interactions, which show fringes that change in size and number as a function of crystal thickness. It has long been known that crystal thickness can be measured to an accuracy of a nm or better using these features in CBED patterns. (Kelly *et al.*, 1975; Allen, 1981) The fine structure of strongly dynamical rocking curves thus gives another way to measure crystal thickness, which should match the minimum $R_1$ for all
reflections. These fringes can be seen clearly for the 311 reflection in Fig. 6. However, the features in Fig. 6f are noticeably sharper than the experimental rocking curve Fig. 6b and this is due to the angular range of the incident electron beam. Additionally, in the optimised orientation Fig. 8b the yellow dots do not lie precisely on a straight horizontal line. Both of these effects may be explained if the intersections are not points, but have a finite size.

The incident electron beam is not a perfect plane wave because the crossover produced by the final condenser lens, which acts as an effective illumination source, is of finite size. We may approximate the angular profile of the incident beam by the intensity profile of the direct transmitted beam averaged through all frames, which is shown in Fig. 10a together with a fit to a Lorentzian profile. The fit is excellent and gives the FWHM of the direct beam to be 0.037 Å⁻¹, or 0.47 mrad (97 arc sec). Thus, the integrated intensity is obtained from the simulated LACBED patterns not from a single row of pixels, but from multiple rows, each of which contributes in proportion with the beam profile of Fig. 10a. This has the effect of changing the rocking curve profile, particularly when the line of the Bragg condition is more parallel to the beam path, and changing integrated intensities because the intensity of a reflection varies along its Bragg condition. The most straightforward way to perform this correction is to apply the beam profile as a convolution to the simulations, allows rocking curves to be extracted that are a good approximation to experiment shown in Figs. 6g and 6h. Integrated intensities obtained after applying this final correction gives $R_1 = 8.9\%$ ($R_2 = 18.1\%, wR_2 = 25.3\%$, Fig. 10b). The largest scatter from the $R_1 = 0$ line is now found in the highest intensity reflections.
Fig. 10. a) The direct beam profile, obtained by averaging all frames in the Si cRED data and a Lorentzian fit. b) $R_1$ calculation after optimising geometry and correcting for slew rate and beam profile (specimen thickness 185 nm). $R_1 = 8.9\%$. c) $R_1$ for the kinematic model $K$, initial dynamical model $D$, with geometry optimisation $G$, corrections for slew rate $S$ and beam profile $P$, and refinement of Debye-Waller factor $DWF$.

Having applied all relevant corrections to the experimental data and optimised the simulation, we are finally in a position to perform a structural refinement. In silicon there is only one free parameter – the Debye-Waller factor $B$. Fig. 11 shows the variation of $R_1$ with $B$, with a final $R_1$ of 6.8\% ($R_2 = 13.8\%$, $wR_2 = 23.6\%$) and a well-defined best-fit at $B = 0.32\text{Å}^2$. This is rather lower than the X-ray value of $B = 0.54\text{Å}^2$ (Többens et al., 2001) and there is still noticeable scatter in the highest intensity measurements. These differences may be due to difficulties in background subtraction, which result from a limited number of intensity measurements in the $PETS$ output (see section S1.1).
Fig. 11. a) $R_1$ as a function of Debye-Waller factor $B$. b) Optimised $R_1$ calculation for $B = 0.33$, giving a best $R_1 = 6.8\%$ at a specimen thickness of 185 nm.

In summary for this silicon cRED data, we have demonstrated that the poor $R_1$ obtained using a kinematical diffraction model is not found when using a dynamical model with appropriate optimisation and correction factors (Fig. 10c). The final result ($R_1 < 7\%$) is still some way above those typical of X-ray diffraction (Ross et al., 2014), indicating that there is still work to do in data processing or modelling for electron diffraction. The importance of careful correction is very clear from the observation that the improvement in $R_1$ due to refinement of the temperature factor (Fig. 11) is of similar magnitude to improvements that result from optimisation of geometry (Fig. 10c). Nevertheless, dynamical ED intensities are far more sensitive to structure (i.e. atomic coordinates) than temperature factors (Beanland et al., 2021), which is reassuring for structure solution and consistent with the growing number of structures solved by ED. Further improvements may be possible with improved background subtraction, which may be incorrect for high intensity reflections in this particular example due to a lack of data in the rocking curves.
4. Discussion and conclusions

In this work, we have established a protocol for dynamical modelling of fine-sliced cRED data, taking account of the corrections that should be applied in the case of electron diffraction, equivalent to those applied to X-ray data. These corrections rely on the ability to extract rocking curves from experimental data, i.e. having a large number of frames collected at small angular increments. We expect that this approach will become widespread as detector technology continues to improve. Our results show similar improvements to those seen by (Palatinus et al., 2013; Palatinus et al., 2015b; Klar et al., 2021), i.e. dynamical modelling of cRED data has a very significant impact on the quality of fit, reducing $R_1$ by almost 20% in the silicon example chosen here. Nevertheless, this is a particularly simple material with very high perfection and many systematic absences. Equivalent improvements may not be found for more interesting (complex) materials, particularly if they have poorer crystallinity, strong inelastic scattering as seen in organic materials (Latychevskaia & Abrahams, 2019), and are not parallel sided lamellae.

Several improvements can be made from this attempt at dynamical modelling, not least the need for significant computing resources. Each set of 70 simulations calculated $5.6 \times 10^6$ incident beam orientations, producing a $400 \times 400$ pixel LACBED pattern for each of the 962 diffracted beams. The large range of reciprocal space covered in each LACBED pattern, together with knowledge of which frame each reflection was seen experimentally, allowed precise correction of the crystal orientation but the area of reciprocal space covered could be reduced by a factor of $> 40$ if this optimisation were first performed by direct calculation of the positions of the different Bragg conditions. To allow rocking curves to be captured fully, each simulation overlapped with the next, meaning that every incident beam orientation was simulated twice, something easily avoided if a single simulation (e.g. $20 \times 14000$ pixels) is calculated instead. Rocking
curves and integrated intensities could be output directly, rather than extracted from these simulated data using python scripts. In *Felix*, the Bloch wave calculation is optimised by careful choice of the diffracted beams included (Zuo & Weickenmeier, 1995; Chuvilin & Kaiser, 2005), but the time required remains $\propto N^3$, where $N$ is the number of beams (Yang *et al.*, 2017). No attempt was made to optimise this parameter and all simulations were run with $N = 200$ from a beam pool of $\sim 800$ in each simulation. If all such improvements were to be implemented it seems reasonable to expect a full set of $I_{hkl}^{(calc)}$ to be obtained in seconds. This would then allow dynamical refinement of crystal structure in reasonable times.

Improvements in simulation fidelity are also possible. The simple Bloch-wave calculation used here assumes that the surface normal is parallel to the incident beam direction, i.e. it does not properly account for continuity of the electron wave function at the entrance and exit surfaces of a tilted crystal. This is obviously not correct for a specimen tilted by up to $70^\circ$. Furthermore, as the crystal rotates, the thickness of material transited by the electron beam changes. Interestingly, we found that simulated rocking curves for a single thickness gave a good match to experiment across the full dataset (section S1.5), and that this thickness agreed with the minimum $R_1$, obtained from integrated intensities. Incorporating a change in crystal thickness $t$ corresponding to that expected for a parallel-sided slab gave no improvement in comparison with a single thickness for the complete data set, although large corrections should only occur at very high tilts ($t \propto 1/\cos \alpha$). We expect a more correct model would yield further improvements in $R_1$. Finally, some experimental rocking curves suffered from poor background correction (section S1.1). While improved background subtraction may be possible in *PETS2* the presence of diffuse scattering, modulated by strong Kikuchi lines, may still be a significant factor that prevents a good measurement of intensity.
In conclusion, dynamical modelling of cRED data has a significant impact on the quality of fit. Improvements are still required before fit metrics for electron diffraction equal those of X-ray diffraction, but there seems to be no fundamental reason why they cannot be achieved. This is encouraging for future development and application of 3D-ED techniques to structural solution in a wide range of applications.

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Supplemental Material for
Modelling fine-sliced three dimensional electron diffraction data with dynamical Bloch-wave simulations

S1.1. Experimental rocking curves and background subtraction.

Four examples of rocking curves from the silicon cRED data produced by PETS, and subsequent background subtraction, are shown in Fig. S1. The red line is a linear least-squares fit to data points outside the peak. The raw integrated intensity is the sum of the green bars. For some strong reflections, such as the 111 peak, the data points in the rocking curve may not extend sufficiently to allow a good fit to the background, leading to a systematic underestimation of integrated intensity. Although not optimised in this data analysis, it is possible to increase the width of the rocking curve extracted by PETS2 by modifying the mosaicity and rocking curve width parameters. It should thus be possible to extract the data in a way that the problem with insufficient "tails" of the rocking curves is avoided.
Fig. S1. Background subtraction for four rocking curves in the Si cRED data set. The line gives the frame by frame output intensity from PETS and the horizontal red line marks the measured background. Green bars show the intensity after background subtraction.
S1.2. Direct beam intensity.

Fig. S2. The relative direct beam intensity, obtained by cropping the stack of \( n = 1389 \) frames to just the direct beam, producing an average of the beam profile by summing all frames and dividing by \( n \), and then dividing each image in the stack by this average. The stack of frames is then re-sliced to view from the side. Deficits in the direct beam, caused by each Bragg condition that is passed through as the goniometer rotates, are visible as dark lines.
S1.3. Reliability factor $R_1$ calculation and sensitivity to $B$ in the kinematic model.

The usual equation for the reliability factor $R_1$ is

$$ R_1 = \frac{\left(\sum |F_{hkl}^{(obs)}| - |F_{hkl}^{(calc)}|\right)}{\sum |F_{hkl}^{(obs)}|}, \quad (S12) $$

where the sums are taken over all observed reflections. We use an equivalent version here for dynamical refinement, with $|F|$ replaced by $I^{1/2}$, i.e.

$$ R_1 = \frac{\sum \left(\sqrt{I_{hkl}^{(obs)}} - \sqrt{I_{hkl}^{(calc)}}\right)}{\sum \sqrt{I_{hkl}^{(obs)}}^{1/2}}, \quad (S13) $$

The quality of fit can be seen by plotting $F_{hkl}^{(obs)}$ against $F_{hkl}^{(calc)}$ as shown in Fig. S3.

![Fig. S3. $R_1$ calculation as a function of $B$ in the kinematic model. The lowest $R$ is found at $B = 0$.](image)
S1.4. Dynamical $R$-factors as a function of thickness.

Since diffracted intensities change significantly as a function of specimen thickness in a dynamical model, each specimen thickness has a different $R_1$. This is illustrated below in Figs. S4 and S5 for Bloch-wave simulations corresponding to the nominal beam path in Fig. 5.

Fig. S4. $R_1$ calculation as a function of specimen thickness $t$ in the dynamic model, with the nominal beam path given by $PETS$. Orange line is a least-squares linear fit.
Fig. S5. $R_1$ as a function of specimen thickness $t$ in the dynamic model, with an optimised beam path and beam profile convolution.
S1.5. *Dynamical rocking curves as a function of thickness.*

The fine structure of rocking curves obtained from strongly dynamical reflections is very sensitive to specimen thickness, as exemplified here by Si 311. The simulated rocking curves have been convoled with the experimentally measured beam profile.

Fig. S6. Comparison of the experimental Si 311 rocking curve (blue) and simulations at a variety of thicknesses (orange)
S1.6. R-factor as a function of B for optimised simulations.

$R_1$ is sensitive to thermal vibrations of atoms in the dynamical model as shown in the plots of $I_{hkl}^{(obs)}$ against $I_{hkl}^{(calc)}$ for different values of the Debye-Waller factor $B$. The systematic underestimation of strong reflections (S1.1) may lead to an underestimation of $B$.

Fig. S7. $R_1$ calculations as a function of $B$ for an optimised dynamical simulation as shown in Fig. 11. The lowest $R$ is found at $B = 0.33$.