1. Introduction

Numerous compounds are known, whose diffraction patterns require four (or more) integer Miller indexes to be indexed completely. Such compounds are said to have modulated structures, which can be interpreted as perturbations with respect to an average conventional structure. Reflections, which cannot be indexed with the usual three-dimensional unit cell, are called satellites. Fourth vector, namely the modulation vector, is introduced as a combination of reciprocal vectors (see Formalism, Eq. (1)), thus creating special cell set of four vectors, called vector modulo of order 4 in three dimensional space. The first indications that crystal structures may not exhibit normal lattice periodicity were found late in 19th century. Physically it means, that neighboring unit cells are not alike, but the deviation of each unit cell from an average structure can be described by a periodic function with its own period. The presence of such periodicity is proved by the existence of a diffraction pattern itself. The formalism of the description was suggested by P. M. de Wolff in 1974 [1]. Reflections that are predominantly strong are consistent with a regular 3-dimensional reciprocal lattice and are called main reflections, but others, called satellites do not fit the same lattice. On the other hand, while the satellites do not fully match the main lattice, the whole pattern can be described by vector modulo of order 4 in three dimensional space. Hence, it can be imagined that the 3-dimensional diffraction pattern is a projection of a 3+d-dimensional reciprocal lattice, in which the main (R) and the satellite (S) reflections are all regularly situated at the lattice nodes (Fig. 1a). From the properties of the Fourier transform the incommensurate modulated structures are treated as 3+d dimensional and real structures can be depicted as three-dimensional sections $R_5$ of a multispace (Fig. 1b, c). Such 3+d unit cells retain the atomic periodicity with atoms represented as strings, while the three-dimensional structure is not periodic. On the other hand, diffraction methods reflect the average atomic distribution over a structure. Thus, the average three-dimensional structure can be treated as the summation of an electron density over additional dimensions ($x_4$ in the case of 3+1 dimensional structure) and its projection on $R_3$. Obviously, the refinement of this type of structure will result in unreal atomic and/or structural parameters, like atomic displacement parameter(s) (ADP), interatomic distances or coordination polyhedra. If the behaviour of atoms along additional dimensions is described by a periodic function, it means, that exact atomic parameters for each unit cell can be calculated. If the ratio between the components of the modulation vector and cell parameter(s) is rational, a structure is called commensurate and can be described in terms of a superstructure, otherwise—this structure is referred to as incommensurate.

It should be stressed, that a structure remains three-dimensional and additional dimensions reflect any structural peculiarity, which the radiation source used is sensitive to. For neutrons it can be orientation of magnetic moments. For X-

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**POWDER DIFFRACTION OF MODULATED AND COMPOSITE STRUCTURES**

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In the past two decades intensive development of the theory of 3+d dimensional crystallography, experimental techniques and corresponding software makes it possible to perform phase analysis and even structure refinement from powder diffraction data for modulated and composite structures. In the present paper the X-ray powder diffraction from such structures is considered. Two examples of multispace structure refinement of the modulated (Bi$_{2.3}$Sr$_{1.7}$CuO$_{6.23}$) and composite (Sr$_{1.32}$Mn$_{0.83}$Cu$_{0.17}$O$_3$) structures from conventional X-ray source data are presented.
rays only positional, occupancy and ADP modulation of atoms are relevant.

Over the next years after P. M. de Wolff publication more stones were put in the foundation of multidimensional crystallography. Practical application of this method came to life. Two programs most commonly used for this purpose were developed: JANA* by V. Petříček and P. Coppens and REMOS† by A. Yamamoto, followed by other software‡. Late in 1980s, first structures were refined from single crystal data with modulation parameters for heavy atoms only. Later light elements were included in this list. Nevertheless, many compounds are not easily grown as single crystals or the interesting phases present complex polydomain structures. Therefore, the study of a structure by single crystal techniques becomes very complicated, if possible. Rietveld refinement from neutron and/or X-ray data is the important step in the development of the multidimensional approach. X-ray powder diffraction from conventional sources is widely used in scientific studies. The correct interpretation of diffraction patterns is important for phase analysis and successive structure refinement. If unindexed peaks remain on a powder XRD pattern of a newly prepared compound, and other preparation routes produce materials having these peaks with similar relative intensities, the following problem should be considered: do extra lines correspond to an admixture, either the superstructure was not correctly taken into account or could they reflect the modulated or composite structure of a pure phase? If these extra lines correspond to satellite reflections of a modulated structure, then their intensities are relatively low. This is not the case for composite structures. Their diffraction patterns include at least two subsets of main reflections with intensities of one order of magnitude and thus, two sets of cell parameters can be refined. Does such a pattern belong to a two phase sample or the sample contains a single phase with the composite structure? In this review we would like to focus on such problems, using our own experience.

2. Short Notes on Formalism

The overwhelming majority of refined structures are described in four-dimensional space, so we will take it as an example. The theory of 3+d dimensional crystallography is described in details so far in many papers including the latest edition of International Tables for Crystallography [2]. We present here a brief description of the four-dimensional case.

The fourth basic vector \( a_4 \), which is perpendicular to the \( a_1, a_2, a_3 \) basic vectors of the \( R_3 \) direct space, is defined through a projection of corresponding reciprocal vector \( a_4^* \) (called modulation vector \( q \)) in \( R_3^* \) reciprocal space (Fig. 1a):

\[
q = \sum_{i=1}^{3} k_i a_i^* = \alpha a^* + \beta b^* + \gamma c^*
\]

where at least one of the coefficients \( \alpha, \beta \) and \( \gamma \) is irrational in general and possibly one or two are rational coefficients. Then, reciprocal cell vectors in \( R_3^* \) space are defined as: \( a_i^* = a_i^* \) for \( i = 1–3 \) and \( a_4^* = q + e_4 \), and from orthogonality condition:

\[
a'_i = a_i - k_i e_4, \quad k_i = \alpha, \beta, \text{ or } \gamma, \quad i = 1–3 \quad a_4 = e_4
\]
The modulation vector is presented by \( \mathbf{q} = \mathbf{q}_0 + m \mathbf{q} \), where \( \mathbf{q}_0 \) and \( \mathbf{q} \) are irrational and rational parts of a modulation vector. The positions of reflections are given by a diffraction vector:

\[
H = h \mathbf{a}^\ast + k \mathbf{b}^\ast + l \mathbf{b}^\ast + m \mathbf{q}
\]

where \( hklm \) are four Miller indices of a reflection. Main reflections (R) correspond to \( m = 0 \), satellite (S) are others (Fig. 1a).

Functions that describe modulation waves are usually expanded in Fourier series:

\[
f^n = A^n + \sum_{m=1} A^n_{c,m} \sin(2\pi n x_4) + \sum_{n=1} A^n_{c,n} \cos(2\pi n x_4)
\]

where \( f^n \) is the actual \( x, y \) or \( z \) coordinate, occupancy or corresponding ADP, \( A^n \) is the corresponding parameter of the average structure, \( A^n_{c,m} \) and \( A^n_{c,n} \) are modulation parameters.

For some special cases the other functions were used, namely linear or saw-tooth function for displacive modulation [3]:

\[
x = x_0 + 2 \ast \mathbf{u}_n \ast \left( \frac{x_4 - x_4^0}{\Delta} \right)
\]

and the step function for occupancy modulation [4]:

\[
\begin{align*}
\rho(x_4) &= 1 \quad \text{if } x_4 \in \left( x_4^0 - \frac{\Delta}{2} , x_4^0 + \frac{\Delta}{2} \right) \\
\rho(x_4) &= 0 \quad \text{if } x_4 \in \left( x_4^0 - \frac{\Delta}{2} , x_4^0 + \frac{\Delta}{2} \right)
\end{align*}
\]

where \( \mathbf{u}_n \) is the amplitude of modulation, \( \Delta \) is the length of the interval, where the function is defined, and \( x_4^0 \) is the parameter of the function representing the center of the step.

Interatomic distances and coordination polyhedra are the most important information for chemists. Two aspects should be taken into account. First, neighboring atoms in the average structure may have different \( x_i \) coordinates. Thus, the comparison of any atoms including interatomic distance calculation should be performed for the same \( R_5 \) section defined by the phase shift parameter \( t \):

\[
t = x_4 - (\mathbf{q} \cdot \mathbf{r})
\]

where \( \mathbf{q} \) is a modulation vector and \( \mathbf{r} \) is a vector defining the position of an atom in the average structure. The modulated structure itself should be considered as a function of any structural parameter (positions, ADPs, distances) vs. \( t \). Second, two neighboring unit cells on the modulation curves do not correspond to nearest points but differ by some value \( \Delta t = t_2 - t_1 \) (\( t_1 \) and \( t_2 \) at Fig. 1b). If we shift the structure along any cell vector by its length, \( \Delta t \) becomes equal to an appropriate component of the modulation vector.

In the case of a composite structure two or more subsystems can be identified, each having its own set of cell parameters and space group symmetry. Let us consider the case with two subsystems, when both of them have common \( a \) and \( b \) parameters and different \( c_1 \) and \( c_2 \). Each subsystem can be defined from common lattice vectors and modulation vector by interlattice matrices:

\[
\begin{pmatrix}
\mathbf{a}_v^* \\
\mathbf{b}_v^* \\
\mathbf{c}_v^* \\
\mathbf{q}_v
\end{pmatrix} = W_v
\begin{pmatrix}
\mathbf{a}_1^* \\
\mathbf{b}_1^* \\
\mathbf{c}_1^* \\
\mathbf{q}_1
\end{pmatrix}
\]

where \( \mathbf{a}_1^* \), \( \mathbf{b}_1^* \), \( \mathbf{c}_1^* \), \( \mathbf{q}_1 \) are reciprocal cell vectors and modulation vector of the composite structure, \( W_v \) is usually unit and \( \mathbf{a}_v^*, \mathbf{b}_v^*, \mathbf{c}_v^*, \mathbf{q}_v \) are reciprocal vectors of the first subcell, reciprocal cell vectors and modulation vector of the \( v \) subsystem. If \( W^2 \) is equal

\[
W^2 = \begin{pmatrix}
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 0 & 1 \\
0 & 0 & 1 & 0
\end{pmatrix}
\]

then \( \mathbf{c}_2^* = \mathbf{q}_1 \). Indices of a reflection in respect to a certain subsystem are defined by:

\[
\begin{pmatrix}
h_v \\
k_v \\
l_v \\
m_v
\end{pmatrix} = (W^*)^{-1} \begin{pmatrix}
h \\
k \\
l \\
m
\end{pmatrix}
\]

For the example given above the main reflections of the first subsystem have \( hkl0 \) indices, while \( hh0m \) reflections are the main reflections for the second one. Reflections with \( hklm \) indices are satellites for both subsystems. Thus, cell parameters of each subsystem can be refined using conventional approach, if necessary, but satellite reflections still can not be indexed in any subsystem using three-dimensional approach.

Symmetry of the 3+1 dimensional modulated
structures is described by four-dimensional space group [2, 5]. In general, each symmetry operation \( (R, \varepsilon, t) \) in the superspace implies \( (R - 3 \times 3 \) rotational matrix, \( t \) – translational vector of a symmetry operation, \( \varepsilon \) and \( t \) refer to the additional dimension):

1. Rotational and translational parts \( (R | t) \) of the symmetry operation belong to the threedimensional space group of the average structure;
2. Point group operation \( R \) must keep the orientation of the modulation vector invariant. Thus, the value of \( \varepsilon \) is defined by the equation, often called de Wolff equation: \( R\varepsilon = \varepsilon R \) or \( \varepsilon R = R\varepsilon \) (index \( i \) means irrational part of the modulation vector);
3. If two atoms \( v, \eta \) are related by a symmetry operation \( (R | t) \) in the average structure, their modulation parameters are related as follows:

\[
\begin{align*}
u_i(x) &= R\varepsilon_i(x - \tau) \\
p_i(x) &= p\varepsilon_i(x - \tau)
\end{align*}
\]

where \( u(x_i) \) is the modulation function of coordinates, \( p(x_i) \) is the occupancy modulation function. Non-zero translational part \( \tau \) is associated with \( \varepsilon = 1 \).

Equations (11) and (12) should be satisfied if atoms \( v \) and \( \eta \) coincide, that is for atoms in special positions, thus putting restrictions on the form of a modulation function. For example, in the case of one harmonic function either \( \sin \) or \( \cos \) waves (if any) should be used, depending on \( \varepsilon \).

For composite structures the symmetries of both subsystems are defined by:

\[
\begin{align*}
R_i &= W^* R_i (W)^{-1} \\
t_i &= W^* t
\end{align*}
\]

where \( R_i \) is the \( 4 \times 4 \) rotational matrix of a symmetry operation and \( t_i \) is the translational part. The subsystems interact, giving rise to modulations of atoms, which are described in respect to their own bases. In addition to main reflections of undistorted subsystems, then satellite reflections occur. In general, such a structure can be described in one subsystem like a modulated structure, but in this case modulations of atoms from the second subsystem will be too large and nonharmonic. A detailed description of a composite structure approach is given in reviews (e.g. S. van Smaalen [6]).

### 3. Powder X-ray Diffraction Patterns

Powder XRD patterns of incommensurate modulated phases have several specific features, which have to be taken into account for successful extraction of information on the unit cell of the average structure and components of modulation vector. The \( hklm \) reflections on powder XRD patterns of incommensurate modulated phase form a dense set being projected on the \( 2\theta \) axis, whereas for non-modulated or commensurate crystals this set becomes discrete. A profile decomposition and Rietveld refinement in the incommensurate case becomes possible because in most cases the intensities of satellite reflections quickly vanish with increasing order (i.e. with increasing \( l m l \) that allows to consider the set of reflections as finite. In the majority of cases only first order satellites are visible on conventional powder XRD patterns. The restriction of maximal value of \( l m l \) allows to assign indexes for satellite reflections. On the other hand, frequent overlapping of weak satellites with intense main reflections and generally low peak to background noise ratio for satellite reflections strongly restrict the accuracy in the positions of the satellites that can be determined from the experimental profile. Unfortunately, poor quality data are usually available from conventional powder XRD pattern for determination and subsequent refinement of the components of the modulation vector. Up to now there is no reliable method, which can help to distinguish a modulated structure and determine a modulation vector from powder data. “...Powder indexing works beautifully on good data, but with poor data it will usually not work at all.” (“Data accuracy for powder indexing” – R. Shirley – NBS Spec. Publ. 567 (1980)), and, in agreement with this statement, a knowledge on three-dimensional reciprocal lattice is still needed for unambiguous determination of length and orientation of the modulation vector in reciprocal space. Selected area electron diffraction in a transmission electron microscope appears to be particularly useful for this purpose since it allows one to observe reciprocal space sections produced by diffraction from small areas of tens nanometers, i.e. from single crystallites from which the powder sample is formed. From the set of such electron diffraction patterns with known mutual orientations a full reconstruction of the three-dimensional reciprocal lattice can be performed. Let us consider several examples of indexing of such patterns. In Fig. 2 the [010] electron diffraction pattern of \( \text{Bi}_{2.3}\text{Sr}_{1.7}\text{CuO}_6.23 \) is shown. The rectangle marks the reciprocal unit cell corresponding to the average structure with the
monoclinic $A2/a$ space symmetry. The satellite reflections form rows assigned to each basic spot; the satellites can be indexed with the modulation vector $q = 0.22a^* + 0.61c^*$. The $\alpha$ and $\gamma$ components of the modulation vector were used then for indexing of powder XRD pattern based on peak positions calculated according to the formula (the monoclinic angle $\beta$ is equal to 90°):

$$1/d_{hklm}^2 = (h + m\alpha)^2/a^2 + k^2/b^2 + (l + m\gamma)^2/c^2$$

Cell parameters and components of the modulation vector could be refined with standard least-squares procedure, as it is done, for example, in the CSD package [7]. For the Bi$_{2.3}$Sr$_{1.7}$CuO$_{6.23}$ phase it gives $q = 0.217(1)a^* + 0.608(1)c^*$, in a good agreement with the electron diffraction data. The indexation of the XRD pattern for Bi$_{2.3}$Sr$_{1.7}$CuO$_{6.23}$ (PDF 45-0315) with four $hklm$ indexes is present in Table 1.

For composite structures two sets of cell parameters should be determined, each belongs to its own subsystem. A separation of reflections belonging to the two main sets becomes easier if the XRD or electron diffraction patterns of both incommensurate and commensurate composite structures are available. A comparison of electron diffraction patterns of Sr$_{4/3}$Mn$_{2/3}$Cu$_{1/3}$O$_3$ and Sr$_{1.32}$Mn$_{0.83}$Cu$_{0.17}$O$_3$ compounds is shown in Fig. 3. This structures consist of two modulated subsystems, namely [(Mn, Cu)O$_3$] and [Sr], which have trigonal lattices with common parameter $a = 9.6$ Å and different parameters $c_1 = 2.6$ Å and $c_2 = 3.9$ Å. The modulation vector can be chosen as $q = c_1/c_2c^*$. The structure of Sr$_{4/3}$Mn$_{2/3}$Cu$_{1/3}$O$_3$ is commensurate (that is $q = 2/3c^*$), then it can be considered as a superstructure with $c = 3 \times 2.6$ Å = 2.90 Å = 7.80 Å. For the Sr$_{1.32}$Mn$_{0.83}$Cu$_{0.17}$O$_3$ the $c_1/c_2 = 0.662$ value requires a large denominator to be approximated by a rational number, and the structure is con-

**Table 1.** X-ray powder diffraction pattern of Bi$_{2.3}$Sr$_{1.7}$CuO$_{6.23}$ (PDF 45-0315). The deviations $\Delta\theta = \theta_{exp} - \theta_{calc}$ do not exceed 0.05°.

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<th>$d$</th>
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sidered as incommensurate. Deviation from being commensurate causes clear splitting of the satellite hklm reflections on the [120] electron diffraction pattern of Sr1.32Mn0.83Cu0.17O3 phase, while the hk0l (from the [(Mn, Cu)O3] subsystem) and hkl0 reflections (from the [Sr] subsystem) remain unsplit on the [120] patterns of both Sr4/3Mn2/3Cu1/3O3 and Sr1.32Mn0.83Cu0.17O3 phases. The indexation scheme of the electron diffraction pattern of Sr1.32Mn0.83Cu0.17O3 phase is shown in Fig. 4, the indexation of powder XRD patterns is given in Table 2. This indexation can be compared with that of the commensurate Sr4/3Mn2/3Cu1/3O3 phase (a=9.5928(8) Å, c=7.8167(6) Å, S.G. P321, Table 3). The details on the Sr1.32Mn0.83Cu0.17O3 structure are described below.

After indexation the set of hklm reflections can be analyzed for the presence of systematic extinctions, and a (3+1)d superspace symmetry group (or several possible symmetry groups) can be proposed based on results of this analysis. This step-by-step procedure in the (3+1)d case is given in International Tables for Crystallography [2] together with a list of (3+1)d superspace symmetry groups.

The correctness of the unit cell, modulation vector and superspace symmetry can be verified by the structure refinement. As well as for
the Rietveld refinement of conventional structures, the initial model for modulated structure comprising atomic coordinates of the average structure and the starting parameters of modulation waves should be defined prior to the refinement. The initial model can be taken from a modulated structure of a similar compound, which was determined earlier from single crystal data, or derived from general structural and chemical considerations, which also can impose several sensible restrictions on the modulated functions thus decreasing a number of refineable parameters. Two such examples of Rietveld refinement of modulated structures are discussed below.

4. Incommensurate Modulated Superconducting Bismuth and Copper Mixed Oxides

Bismuth and copper mixed oxides were first synthesized by B. Raveau with co-workers [8]. Some of these substances were observed to exhibit superconducting properties, which resulted in extensive study of these compounds. The problem of the structure determination and refinement occurred at once due to significant modulations in their structures.

Bi-containing layered copper oxides Bi$_{2-x}$Sr$_x$Ca$_2$Cu$_{2n}$O$_{2n+4}$ (n=1–3) have an intergrowth structure with perovskite and rock salt fragments alternating along the c axis (Fig. 5). They have an incommensurate modulated structure, and the appearance of modulation is associated with a mismatch between the BiO layer and perovskite-like fragment. This mismatch is reduced via strong shifts of atoms in all layers from ideal sites and insertion of additional oxygen atoms in the bismuth-containing layer. This model was suggested by H. Zandbergen et al. [9] and later proved by numerous studies.

An influence of modulations on a distribution of scattering density in Bi-2201 (n=1) phases is illustrated by a comparison of two high-resolution electron microscopy images. These

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**Table 2.** X-ray powder diffraction pattern of Sr$_{1.32}$Mn$_{0.83}$Cu$_{0.17}$O$_3$. The deviations $\Delta \theta = \theta_{exp} - \theta_{calc}$ do not exceed 0.05°. Reflections with indices in two subsets are common for both subsystems. Only reflections with $\|I/I_{100}\| \geq 1\%$ are shown.

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**Table 3.** X-ray powder diffraction pattern of Sr$_{4/3}$Mn$_{2/3}$Cu$_{1/3}$O$_3$. The deviations $\Delta \theta = \theta_{exp} - \theta_{calc}$ do not exceed 0.05°. Only reflections with $\|I/I_{100}\| \geq 1\%$ are shown.

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images show the BiO layers in the modulated Bi$_2$Sr$_{1.6}$La$_{0.4}$CuO$_6.33$ phase (Fig. 6, top) and in the same phase after strong fluorination, where the modulations are suppressed by fluorine insertion into the BiO slabs (Fig. 6, bottom) [10].

Apparently, the structure refinement of Bi-2201 phase without modulations should result in a rough model with very large atomic displacement parameters especially for Bi atoms and unrealistic long Bi–O distances. The location of extra oxygen cannot be revealed within such model. The refinement of incommensurate modulated structures was successfully made from single crystal data [11–13]. However, it is very difficult to grow single crystals of Bi containing cuprates with desired composition. Several studies were performed using powerful sources such as neutron TOF or/and synchrotron radiation to obtain powder data and structures were refined [14, 15]. Although satellite reflections are much weaker than main reflections, their maximum intensities may be as high as 5–10% (relative scale). The significance is, that structure refinement from powder data without taking modulations into account can result in poor agreement factors. Now we consider the refinement of modulated bismuth cuprates from powder data and compare results with single crystal refinement. Sample preparations and data collections are described previously [16, 17]. The structures were refined using JANA2000 program package [18, 19].

A powder sample with the composition Bi$_{2.3}$Sr$_{1.7}$CuO$_6.23$ was used in this study. The 2201 phases crystallize in $A2/(a0\gamma)$ monoclinic superspace group with pseudotetragonal cell parameters: $a=5.4$ Å, $c=24.5$ Å. First, the average structure was refined with the initial parameters taken from Leligny et al. [20]. Strontium site was supposed to be occupied by Sr and Bi with
a 0.85:0.15 ratio. The refinement converged only with a very low damping factor, fixed x coordinates for Bi and Sr atoms, and resulted to poor agreement with the experimental and calculated data ($R_f=0.096, R_p=0.112$). Such result was expected, because there are about 20–25 satellite reflections below 2\(\theta\)=50° (for CuK\(\alpha_1\) radiation), which can be detected by profile decomposition of the pattern.

In incommensurate structure refinements several reliability factors are calculated according to the separation of the reflections into the group of main reflections and the groups of satellites of different order. $R_f$ or $R_p$ factors should be defined for each set of reflections. In general, satellite peaks are characterized by low peak to background ratios. This characterization leads to a significant effect on integral intensities of the satellites by the background noise and to larger reliability factors for these sets of reflections, even if the structure model is correct. Only first order satellites were used in the refinement of modulation parameters, higher order satellites have too low an intensity in the powder pattern and their intensities were not detected with profile analysis.

Displacive modulation parameters (one harmonic) for Bi atom were introduced in the refinement, resulting in a significant reduction of residuals: $R_{F0}=0.072, R_{Fm}=0.052, R_{F1}=0.088, R_p=0.087$ (here and further on index 0 stands for overall, $m$ – for main reflections, $1$ – for first order satellite reflections). Starting parameters for modulation were also taken from Leligny et al. [20]. Modulation parameters for other cations were then refined. Introduction of one harmonic for displacive modulation of all cations resulted in $R_{F0}=0.049, R_{Fm}=0.036, R_{F1}=0.060, R_p=0.073$. The refinement of the second harmonic revealed significance over 3\(\sigma\) only for Bi modulation parameters and along $x_3$ for Sr. At this stage $R$-factors were reduced to $R_{F0}=0.039, R_{Fm}=0.027, R_{F1}=0.049, R_p=0.068$. ADPs for all oxygen atoms were fixed to $U_{iso}=0.025$ and were not refined. For oxygen modulation refinement the model suggested in [21] was used. Modulations of oxygen atom in CuO\(_2\) layer were refined with a harmonic function only. For other oxygen atoms (in the BiO and SrO layers) more complicated scheme was chosen. Modulations along $x_1$ were refined by a linear function with $\Delta=1$, while along $x_2$ and $x_3$ were left harmonic. Additional oxygen atom in BiO layer was not included into the refinement since its impact in powder XRD pattern is too small to be properly taken into account. The refinement converged only with low damping factor and reduced $R$-factors to $R_{F0}=0.036, R_{Fm}=0.026, R_{F1}=0.045, R_p=0.065$. Experimental, calculated and difference powder patterns are presented at Fig. 7.

The results of the refinement of the modulation functions for heavy scattering cations are very similar to those obtained from single crystal data for Bi\(_{2.26}\)Sr\(_{1.74}\)CuO\(_{6.19}\) [21]. Amplitude of

![Fig. 7. Experimental, calculated and difference powder XRD patterns for Bi\(_{2.3}\)Sr\(_{1.7}\)CuO\(_{6.23}\). Ticks mark the position of the main hkl0 reflections.](image)
modulations and t regions with maximal shift are essentially the same, some examples are presented at Fig. 8. As a result we may state, that the refinement of modulated structure from X-ray powder data produces a rough model of structural modulation. It reflects the general tendency and may be used for indexing of powder data, but exact atomic parameters and distances are the rough approximation of the real structure. The importance of such refinement is connected with full indexing of a diffraction pattern. The situation can be significantly improved if structural constrains are used in the refinement. In comparison with a single crystal X-ray diffraction experiment, X-ray powder patterns provide only information on a much smaller amount of structure amplitudes. Therefore, the number of refineable structure parameters should be restricted to those, which are very essential for the structure description. Now we will consider such approach for the composite Sr\textsubscript{1.32}Mn\textsubscript{0.83}Cu\textsubscript{0.17}O\textsubscript{3} structure.

5. Sr\textsubscript{1.32}(Mn, Cu)O\textsubscript{3} Hexagonal Perovskite Based Oxides with Chain Structures

Sr\textsubscript{1.32}Mn\textsubscript{0.83}Cu\textsubscript{0.17}O\textsubscript{3} belongs to the family of hexagonal perovskite based oxides with the common formula A\textsubscript{2m+n},A'\textsubscript{n},B\textsubscript{3m+n},O\textsubscript{9m+6n}. The structure of these oxides is generally considered as a derivative of the 2H hexagonal perovskite ABO\textsubscript{3} (Fig. 9a) that is based on the (hh) close packing of the (A\textsubscript{3}O\textsubscript{9}) layers. This structure consists of infinite chains of face-shared BO\textsubscript{6} octahedra with columns of A cations in between forming a hexagonal array in the a=apar/2, c=(2/√3)apar unit cell. One third of the oxygen atoms can be removed from the (A\textsubscript{3}O\textsubscript{9}) layers in an ordered manner, so that one trigonal prismatic cavity is created instead of two face sharing octahedral polyhedra (Fig. 9b). Additional A' cations can be located inside these trigonal prisms resulting in a (A\textsubscript{3}A'O\textsubscript{6}) layer composition. The ordered alternation of m (A\textsubscript{3}O\textsubscript{9}) and n (A\textsubscript{3}A'O\textsubscript{6}) layers leads to numerous A\textsubscript{2m+n},A'\textsubscript{n},B\textsubscript{3m+n},O\textsubscript{9m+6n} polytypes described by J. M. Perez-Mato et al. [22]. A comprehensive list of the A\textsubscript{2m+n},A'\textsubscript{n},B\textsubscript{3m+n},O\textsubscript{9m+6n} compounds is given in [23]. The different stacking sequences of the (A\textsubscript{3}O\textsubscript{9}) and (A\textsubscript{3}A'O\textsubscript{6}) layers results in different values of c parameter and often leads to an enormous increase, which makes the structure refinement impossible. The composite structure model was first suggested by M. Evain et al. [24] and generalized by J. M. Perez-Mato et al. [22]. These structures are con-
sidered as consisting of \([(B, A')O_3]/H11009\) and \([A]/H11009\) subsystems with a common parameter, difference \(c_1\) and \(c_2\) parameters and a modulation vector defined as \(\mathbf{q} = \gamma \mathbf{c}^*\). In general they are commensurate with \(\gamma = 3(n + m)/2(2n + 3m)\), but for members with large denominators it is better to consider the modulation as incommensurate.

Intensities of satellite reflections for modulated structure are rather weak and sometimes could be ignored. Composite structures provide reflections from both sublattices with intensities of the same order of magnitude that should definitely taken into account. Commensurate structures can be treated as the superstructure, while in the incommensurate case \(\Delta d\) approach is the only way out.

The synthesis and data collection for \(\text{Sr}_{1.32}\text{Mn}_{0.83}\text{Cu}_{0.17}\text{O}_3\) are described elsewhere [25]. The crystal structure of \(\text{Sr}_{1.32}\text{Mn}_{0.83}\text{Cu}_{0.17}\text{O}_3\) was refined from X-ray powder diffraction data in \(R\bar{3}m\) (00l) superspace group with two sets of cell parameters obtained from powder pattern profile analysis and successive least-squares refinement: \(a_1 = 9.5971(8)\ \text{Å}, \ c_1 = 2.60008(17)\ \text{Å}, \ c_2 = 3.9312(3)\ \text{Å}\), thus producing modulation vector \(\mathbf{q} = 0.6616(1)c^*\) with interlattice matrices:

\[
W^1 = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}, \quad W^2 = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{pmatrix}
\]

The incommensurate composite model for the \(A_{x+y}A'B_{x+y}O_{6+y}\) compounds has been previously discussed, and will be briefly summarized here. The average structure is presented as \([\text{Mn}_{0.83}\text{Cu}_{0.17}\text{O}_3]_n\) \((\text{Mn, Cu site} = 0, 0, 0; \ \text{O site}^* = 0.16, 0.16, 1/2)\) in the first subcell and \([\text{Sr}]_n\) \((\text{Sr site} = 1/3, 0, 1/4)\) in the second one. The Sr content is determined by a volume ratio \(V_1/V_2 = 0.662\), where \(V_1\) and \(V_2\) are cell volumes of the first and second subsystems, respectively. The amount of the \(A\) cations determines \(g = (1 + x)/2\), where \(x\) is taken from the \(A_{1-x}(A', B_{1-x})\text{O}_3\) formula. The occupancy modulations for Sr, \((\text{Mn, Cu})\) and O atoms are modeled with the saw-tooth functions.

The saw-tooth function with \(\Delta = 1/2\) and \(x_0^2 = 1/4\) for oxygen atoms with the average coordinates \((0.16, 0.16, 1/2)\) separates them into two sets of atoms related by a mirror plane \(m_{010}\) and \(Ob\), so that each set forms the \((001)\) triangular face of an octahedron or a trigonal prism. Depending upon the \(t\) coordinate, the trigonal prismatic sites are created by either \(Oa–Oa\) or \(Ob–Ob\) combinations, whereas the \(Oa–Ob\) combination corresponds to an octahedral site (Fig. 10, top). In the idealized structure model of \(A_{3m}A'B_{3m}O_{9m+6n}\) hexagonal perovskites the width of the octahedron \(D_O\) measured along the \(c\) axis is one half the width of the trigonal prism \(D_p\), however in reality the \(D_p = 2D_O\) condition is not fulfilled. To overcome this difference a displacement of oxygen atoms along the \(c\) axis is introduced, which is characterized by the \(u_{0,3}(O) = (D_p – D_O)/2\) component \(u_{0,1}(O)\) and \(u_{0,2}(O)\) were assumed to be small and fixed to zero). The \((\text{Mn, Cu})\) cations at \((0, 0, 0)\) are separated into two sites, one situated at the B octahedral sites and the other situated at the \(A'\)

* The site is half occupied; see the further explanation.
trigonal prismatic sites. Windows of the saw-tooth functions for B and A' positions should coincide with the regions where either octahedral or prismatic oxygen environment occurs, that requires \( \Delta_\text{A} = (1-x)/2 \) and \( \Delta_\text{B} = x/2 \) and \( x^2_{\text{B}} = 0 \), \( x^2_{\text{A}} = 1/4 \) (Fig. 10, bottom). By symmetry \( u_{\text{B}}(\text{B}, \text{A}') = u_{\text{B}}(\text{A}', \text{B}) = 0 \), whereas \( u_{\text{B}}(\text{B}) \) and \( u_{\text{B}}(\text{A}') \) should result in the same slope of the saw-tooth functions as for oxygen atoms. Hence, \( u_{\text{B}}(\text{B}) = (1-x)u_{\text{B}}(\text{O}) \) and \( u_{\text{B}}(\text{A}') = xu_{\text{B}}(\text{O}) \). Thus, the distribution of the cations between octahedral and prismatic sites is determined by a single refineable parameter \( u_{\text{B}}(\text{O}) \).

In the second subsystem, the Sr atoms at \((1/3, 0, 1/4)\) can be located along the \( c \) axis either at the level of the centers of the prisms or at the level of the triangular faces. These two subsets are described by saw-tooth functions with \( \Delta_{\text{Sr}1} = x(1+x), x^2_{\text{Sr}1} = 0 \) and \( \Delta_{\text{Sr}2} = (1/3) - [x(1+x)], x^2_{\text{Sr}2} = 1/2, u_{\text{Sr}}(\text{Sr}1) = [3x(1-2x)]u_{\text{Sr}}(\text{Sr}2) \).

This model was introduced to the structure refinement taking into account satellite reflections up to second order. The preferred orientation was refined with the March-Dollase function. The initial starting model gave reasonable values of reliability factors \( R_{\text{all}} = 0.057, R_p = 0.017 \). Due to the small difference in scattering power of Mn and Cu we did not refine the occupancy factors for the B and A' sites to determine the distribution of these cations. The B site was assumed to be fully occupied by Mn, whereas 53.9% Cu and 46.1% Mn were placed at the A' site to achieve the nominal phase composition. After the initial refinement step large value of atomic displacement parameter for the A' cation \((U_{\text{iso}} = 0.024 \text{Å}^2)\) was observed. The A' cations were displaced from their ideal position \((0, 0, 0)\) to a \((x, x, 0)\) position, that means a shift of these cations towards one of the rectangular faces of the trigonal prism. This shift results in a two times decrease of the atomic displacement parameter for these atoms, followed by a decrease of the reliability factor to \( R_{\text{all}} = 0.049 \). Further improvement of the fit was achieved by adding one harmonic component of the positional modulation function to the saw-tooth displacement of the B cations. Prior to that, the orthogonalization procedure was performed for the modulation functions of the B cations [26]. The final values of the reliability factors \( R_{\text{all}} = 0.044, R_p = 0.016 \) reflect the good agreement between calculated and experimental profiles (Fig. 11).

Let us consider the chemical formula \( A_{1-x}A'_{x} \) \( \text{B}_{1-x} \text{O}_3 \), where \( x = p/q \), where \( p \) and \( q \) are co-prime numbers. If \( N_{\text{prism}} \) and \( N_{\text{oct}} \) are respectively the numbers of trigonal prismatic and octahedral cavities in \( \{\text{B}, \text{A'}\}\text{O}_3 \) chains, then the ratio \( N_{\text{prism}}/N_{\text{oct}} = x(1-x) - p(q-p) \). BaNiO\(_3\) [27] \((N_{\text{prism}} = 0, N_{\text{oct}} = -1, \text{ consequently } p_1 = 0, q_1 = 1)\) (Fig. 9a) and Sr\(_4\)PtO\(_6\) [28] \((N_{\text{prism}} = -1, N_{\text{oct}} = -1, \text{ consequently } p_2 = -1, q_2 = 2)\) (Fig. 9b) can be taken as parent structures. The combination of these structures will produce a new structure with

![Fig. 11. Experimental, calculated and difference X-ray diffraction profiles for Sr\(_{1.32}\)Mn\(_{0.83}\)Cu\(_{0.17}\)O\(_3\).](image-url)
$p/q = (p_1 + p_2)/(q_1 + q_2) = 1/3$. This is the structure of the commensurate Sr$_{1.3}$Mn$_{0.3}$Cu$_{1.0}$O$_3$ phase ($N_{\text{prism}} = 1$, $N_{\text{oct}} = 2$, consequently $p_1 = 1$, $q_1 = 3$). The process is continued to infinity, giving rise to enormous number of phases as described by Farey tree [22]. For Sr$_{1.3}$Mn$_{0.3}$Cu$_{1.0}$O$_3$ the refined value of $\gamma = 0.6616(1)$ corresponds to $x = 0.3232$. It is close to $p/q = 1/3$, where $p$ is the number of prisms in the chains and $(q-p)$ is the number of octahedra. Using Farey tree [22], the fractional value $p/q = 32/99 = 0.3232$ for $x$ can be found. Expressing $p/q$ through the preceding “generators” $p_1/q_1$ and $p_2/q_2$, so that $p/q = (p_1 + p_2)/(q_1 + q_2)$ the sequence of octahedra and prisms can be found. Thus, the repeat period along the chains comprises 99 polyhedra according to the sequence 1×(3 oct + 1 prism) + 9×(2 oct + 1 prism) + 1×(3 oct + 1 prism) + 10×(2 oct + 1 prism) + 1×(3 oct + 1 prism) + 10×(2 oct + 1 prism). Such polyhedral arrangement was confirmed by high resolution electron microscopy [25].

6. Concluding Remarks

The intrinsic one-dimensional character of powder XRD pattern results in a significant loss of structural information if compared with the single crystal X-ray diffraction technique. A lack of data seems to be crucial for possibility to determine \textit{ab initio} the structure model for an incommensurate modulated compound using powder XRD data as a unique source of information. However, compared with other techniques such as electron diffraction and high-resolution electron microscopy, powder XRD patterns of incommensurate phases become applicable not only for phase analysis, but also for determination and precise refinement of the unit cell metrics, modulation vector and possible superspace group. If an appropriate model is available from a similar known structure or from chemical considerations, Rietveld refinement appears to be able to provide sensible information on modulation parameters and shed light on peculiarities of the structural organization of composite structures.

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