

## A Standard Numbering Scheme for Class C $\beta$ -Lactamases

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**ABSTRACT** Unlike for classes A and B, a standardized amino acid numbering scheme has not been proposed for the class C (AmpC)  $\beta$ -lactamases, which complicates communication in the field. Here, we propose a scheme developed through a collaborative approach that considers both sequence and structure, preserves traditional numbering of catalytically important residues (Ser<sup>64</sup>, Lys<sup>67</sup>, Tyr<sup>150</sup>, and Lys<sup>315</sup>), is adaptable to new variants or enzymes yet to be discovered and includes a variation for genetic and epidemiological applications.

**KEYWORDS** AmpC, amino acid numbering, beta-lactamases, class C beta-lactamase, conserved residue, nomenclature, structure-activity relationships

An urgent need exists to address current inconsistencies in the numbering of amino acid residues among class C  $\beta$ -lactamases, both within families and across the class. Established conventions in the field define three common features shared among the serine-type  $\beta$ -lactamases. In the class C  $\beta$ -lactamases, also known as AmpC  $\beta$ -lactamases, these features occur at recognizable conserved motifs: S<sup>64</sup>XXK (where S<sup>64</sup> is the active-site serine), Y<sup>150</sup>XN, and K<sup>315</sup>(S/T)G (1–4). These designations align with the amino acid sequence of the mature form of both the P99 AmpC (originally characterized in an *Enterobacter cloacae* strain [NCBI RefSeq accession no. WP\_049134845.1] but now found to be an *Enterobacter hormaechei* strain [GenBank accession no. CAA30257.1]) and *Escherichia coli* AmpC (NCBI RefSeq accession no. WP\_001336292.1; originally characterized in strain K-12 under GenBank accession no. AAC77110.1). While *E. coli* AmpC has historical significance as both the first  $\beta$ -lactamase reported (5) and the first class C  $\beta$ -lactamase sequenced, P99 maintains the same numbering of conserved motifs and the mature form begins with residue one (6) rather than residue four as in *E. coli* AmpC. In contrast, many other class C  $\beta$ -lactamases possess insertions and deletions that shift the numbering of the conserved residues, significantly complicating both nomenclature and comparisons between enzymes.

For this report, we analyzed 155 unique AmpC structures deposited in the Protein Data Bank (including 142 supported by 66 publications) and found that 129  $\beta$ -lactamase structures identify the catalytic serine as Ser<sup>64</sup> (123 naturally and 6 with alignment), 10 number from the beginning of the precursor form with the signal peptide included, and the remaining 16 number from the beginning of the mature form but do not identify the catalytic serine as Ser<sup>64</sup> (of which 8 are not associated with a publication). Additionally, based on a literature search of PubMed, we found that consistency is lacking for numbering within the various families of class C  $\beta$ -lactamases. As an example, since the term PDC (*Pseudomonas*-derived cephalosporinase) was coined in 2009 for the chromosomal AmpC of *Pseudomonas aeruginosa*, three different approaches have been used to number amino acid residues in this  $\beta$ -lactamase (7). These approaches include (i) direct numbering of residues beginning with the N terminus of the precursor protein (7), (ii) direct numbering of residues beginning with the N terminus of the mature protein (8), and (iii) alignment-based numbering designed to maintain the conventional assignment of conserved residues and to simplify numbering for comparisons across families (9). Unfortunately, it can be unclear to readers which of the various schemes is being used in a given publication. As a result, authors may sometimes find choosing a numbering scheme and numerically designating a given residue problematic. Comparing findings from multiple publications may be made unnecessarily difficult; resolving ambiguity in assignment may be extremely challenging. For example, a reference to Gly at position 183 in PDC may refer to a site that is described as having a clinically relevant mutation if numbering begins with Met<sup>1</sup> of the precursor form but would refer to a different glycine, 26 residues away, if alignment-based numbering was used (10, 11).

To address this growing concern, we propose a numbering scheme to use consistently when referring to crystallographically equivalent positions in the mature form of any class C  $\beta$ -lactamase. We suggest the acronym “SANC” to name the scheme, for structural alignment-based numbering of class C  $\beta$ -lactamases, or else the simpler term “structural position.” In developing this numbering scheme, we adapted the approaches used by Ambler et al. for the class A  $\beta$ -lactamases (12) and Galleni et al. for the class B  $\beta$ -lactamases (13). We conducted an amino acid alignment of 32 AmpC  $\beta$ -lactamases, both chromosomal and plasmid encoded (see Table S1 in the supplemental material) and identified characteristic differences from P99 for each enzyme (Table 1). Sequences were obtained from the National Center for Biotechnology Information Protein Database (14), and signal peptide cleavage

**TABLE 1** Insertions and deletions present in the AmpC enzymes examined in comparison to *E. cloacae* complex P99

Class C $\beta$ -lactamase	NCBI accession no.	Insertions and deletions relative to <i>E. cloacae</i> complex P99 <sup>a</sup>
ACC-1	WP_032491956.1	-116, +204a, +247a, -289, -290, +362, +363
ACT-1	WP_063857727.1	-361
ADC-7	WP_063857816.1	+0, +204a, -245, -304, -305, -306, +362
ADC-8	WP_004923134.1	+0d, +0c, +0b, +0a, +0, -245, +362, +363, +364, +365, +366, +367
AQU-1	WP_099156042.1	-1, -2, +204a, -243, -245, -301, -302, +362
<i>Burkholderia multivorans</i> AmpC1	WP_012218336.1	+204a, -245, +362, +363, +364
BUT-1	WP_104531863.1	+0a, +0
CepH	WP_063843234.1	-1, -2, +204a, -243, -245, +362
CepS	WP_063843235.1	-1, -2, +204a, -243, -245, +362
CFE-1	WP_032490699.1	None
CMA-1	WP_032974004.1	-1, -2, -3, -4, -5, -6, -116, +204a, -245, +362
CMH-1	WP_063859580.1	None
CMY-2	WP_000976514.1	None
CSA-1	WP_007888761.1	-1, -2, -3, -4, -5, -6, -116, +204a, -245, +362
DHA-1	WP_004236386.1	-1, -2, -3, -4, -301
<i>Escherichia coli</i> AmpC	WP_001336292.1	-1, -2, -3
EC-5	WP_001443153.1	-1, -2, -3
<i>Edwardsiella</i> AmpC	WP_041692555.1	+0a, +0
FOX-4	WP_032489727.1	-1, -2, +204a, -243, -245, +362
LHK-1	WP_081666691.1	-1, -2, -3, -4, +204a, -245
LRA-10	WP_099982803.1	-1, -126, +204a, -245, -361
LRA-18	WP_099982801.1	-1, -245, -311, +362, +363, +364, +365
<i>Mycobacterium smegmatis</i> AmpC	WP_011729443.1	-1, -2, -3, -4, -5, -6, +204a, -245, -305, -306, +362
MIR-1	WP_032489464.1	None
MOX-1	WP_032489888.1	+0, +204a, -243, -245, -301, -302, -303, +362
OCH-1	WP_040129485.1	+0, +204a, -245, +362, +363, +364
PAC-1	WP_034051940.1	-1, -2, -3, -4, -5, -116, +204a, -245, +362
PDC-1	WP_003101289.1	+125a, +204a, -245, +362, +363, +364, +365, +366, +367, +368, +369, +370
SRT-1	WP_063864749.1	-1, -2, -3, -4, -5, -6, -116, +204a, -245, +362, +363
SST-1	WP_063864750.1	-1, -2, -3, -4, -5, -6, -116, +204a, -245, +362, +363
TRU-1	WP_042027926.1	-1, -2, +204a, -243, -245, +362

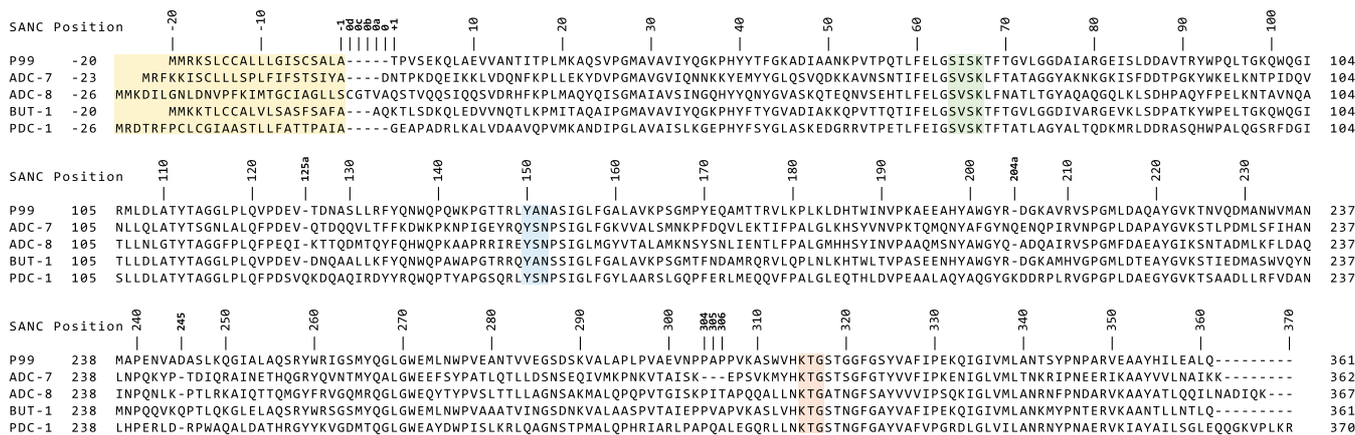
<sup>a</sup>A minus indicates a deletion, and a plus indicates an insertion. Appended letters indicate that an insertion follows a given residue number.

sites were determined using UniProt (or SignalP 5.0 for entries not present in UniProt) (15, 16). Mature protein sequences were aligned using the MUSCLE algorithm (17) with default settings.

Consensus secondary structure (defined as a majority of structures in agreement for a given amino acid position) was determined based on comparisons of a representative structure of each of the 10 AmpC  $\beta$ -lactamases for which one or more structures are available in the Protein Data Bank, specifically, ACT-1 (PDB ID [2ZC7](#)), ADC-7 (PDB ID [4U0T](#)), CMY-2 (PDB ID [1ZC2](#)), *E. coli* AmpC (PDB ID [2BLS](#)), FOX-4 (PDB ID [5CGS](#)), MOX-1 (PDB ID [3W8K](#)), *Mycobacterium smegmatis* AmpC (PDB ID [5E2H](#)), PDC-1 (PDB ID [4GZB](#)), and TRU-1 (PDB ID [6FM6](#)). The consensus agrees with the secondary structure (or lack thereof) of P99 for just over 90% of residues. This consensus was used to annotate secondary structure, including stripes to indicate residues with an even split between two secondary structure types, and helix numbers on the alignment. Finally, a simple literature survey was conducted to determine residues belonging in either the consensus portion or the fullest likely extent of the  $\Omega$ -loop or R2-loop, both of which are also annotated on the alignment. By including this structural information, we hope to both better correlate the numbering system with well-known structural features and to provide additional points of reference for those just beginning to work with AmpC structures.

The exact positions of one insertion and one deletion within the alignment were manually adjusted (residue 203a by MUSCLE became 204a by structure to preserve a  $\beta$ -turn, and the deletion of residue 247 by MUSCLE became a deletion of residue 245 by structure to preserve an  $\alpha$ -helix) to ensure that they occurred in structurally reasonable areas of both the consensus structure and 10 source structures.

Amino acid numbering was based on *E. cloacae* complex P99 while preserving the conventional numbering of the following residues: Ser<sup>64</sup>, Lys<sup>67</sup>, Tyr<sup>150</sup>, and



**FIG 1** Use of alignment to assign SANC-based amino acid residue numbers. Positions corresponding to insertions and deletions are indicated in bold. ADC-7 adds residues 0, 204a, and 262 and deletes residues 245 and 304 to 306. ADC-8 adds residues 0 to 0d and 262 to 267 and deletes residue 245. BUT-1 adds residue 0. PDC-1 adds residues 125a, 204a, and 362 to 370 and deletes residue 245. For reference, signal sequences are highlighted in yellow, S<sup>64</sup>XXX in green, Y<sup>150</sup>XN in blue, and K<sup>315</sup>(S/T)G in red.

Lys<sup>315</sup>. Insertions relative to P99 were addressed by appending a lowercase letter(s) to the number of the amino acid immediately preceding the insertion (e.g., 125a in PDC-1). Deletions relative to P99 were skipped, resulting in “ghost residues” (e.g., ACC-1 has residues G115 and L117 with a deleted residue at position 116). For mature enzymes with more C-terminal amino acid residues than P99, additional residues are assigned numbers in numerical order beginning with 362. For mature enzymes with more N-terminal amino acid residues than P99, the first additional residue is numbered 0 and subsequent residues are numbered by appending a lowercase letter to zero while moving in an N-terminal direction (e.g., 0 and 0a for BUT-1 and *Edwardsiella* AmpC). Signal peptide residues are assigned negative numbers, beginning with -1 for the residue adjacent to the cleavage site and proceeding in the N-terminal direction until all residues are numbered. Multiple sequence alignments are not considered for the signal peptide regions. Figure 1 illustrates these principles with several examples.

Amino acid positions should be provided under both a family-specific, precursor-based scheme (precursor numbering) and the alignment-based scheme (SANC) at first mention of a given residue in a publication. Authors are free to choose their favored convention for subsequent mentions, but as a general suggestion, we encourage the use of SANC for biochemical and structural publications and precursor numbering for genetic and epidemiological publications.

Providing numbering under both schemes is essential to our proposal. Structural numbering maintains continuity with the conventional assignment of the catalytic serine as Ser<sup>64</sup> and the majority of existing literature on class C  $\beta$ -lactamase structure and function, while precursor numbering enables direct gene translation and simplifies interpretation of sequencing results, particularly within a single family. Utilizing this hybrid approach, an initial description of a typical PDC variant might read “PDC-221 differs from PDC-1 (GenBank accession no. [AAG07497.1](http://www.ncbi.nlm.nih.gov/nuccore/AAG07497.1)) by a single amino acid substitution, E247K, occurring at SANC position 219.”

In Table S1 in the supplemental material, we show a multiple sequence alignment of 32 class C  $\beta$ -lactamases with column headers indicating the appropriate number to be used at each position. The spreadsheet also features a text-based alignment of the structures used in determining the consensus secondary structure. In Text S2 in the supplemental material, we provide a protein profile hidden Markov model (HMM) which implements the SANC scheme, built from the multiple sequence alignment using HMMER (<http://hmmer.org>). Alignments of the HMM to class C  $\beta$ -lactamases are expected to produce correct SANC assignments when results of the search are examined. We suggest using the HMM, rather than examinations by eye, to make position

assignments under this scheme for novel AmpC enzymes that may be discovered in the future. Finally, basic instructions for using our HMM with the HMMER software are also included in Text S1 in the supplemental material.

For the specific case of PDC variants, a database utilizing the three numbering schemes (SANC and both family-specific precursor and mature form numbering) is freely available at <https://arpbigidisba.com/pseudomonas-aeruginosa-derived-cephalosporinase-pdc-database/>.

## SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1**, XLSX file, 0.1 MB.

**SUPPLEMENTAL FILE 2**, PDF file, 0.5 MB.

**SUPPLEMENTAL FILE 3**, TXT file, 0.2 MB.

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