A minimum reporting standard for multiple sequence alignments

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Multiple sequence alignments (MSAs) play a pivotal role in studies of molecular sequence data, but nobody has developed a minimum reporting standard (MRS) to quantify the completeness of MSAs. We present an MRS that relies on four simple completeness metrics. The metrics are implemented in AliStat, a program developed to support the MRS. A survey of published MSAs illustrates the benefits and unprecedented transparency offered by the MRS.

MSAs are widely used during annotation and comparison of molecular sequence data, allowing us to identify medically important substitutions\(^1\), infer the evolution of species\(^2\), detect lineage- and site-specific changes in the evolutionary processes\(^3\) and use ancestral sequence reconstruction to engineer new enzymes\(^4\). There is a wide range of computational tools to obtain MSAs, and two of these (i.e., Clustal W\(^5\) and Clustal X\(^6\)) are now among the 100 most cited papers in science\(^7\).

MSAs often have gaps inserted between the nucleotides or amino acids of some of the sequences. These gaps are inserted to maximize the homology of residues from different sequences. A correct MSA is necessary for accurate genome annotation, phylogenetic inference and ancestral sequence reconstruction. However, deciding where to put the alignment gaps may be more art than science. This is because homology is defined as similarity due to historical relationships by descent\(^8\). Most of these relationships belong to the unobservable distant past, so it is impossible to measure the accuracy of most MSAs inferred from real sequence data.

Without this ability, reporting the completeness of MSAs may be the best that can be achieved. So far, the only metric sometimes used is the percent missing data for a sequence\(^9\) or an alignment\(^10\), but neither is sufficiently transparent and insightful. To ameliorate this, we developed a minimum reporting standard (MRS) for MSAs.

The MRS uses four metrics to quantify the completeness of different attributes of MSAs. Given an MSA with \(m\) sequences and \(n\) sites, we may compute four metrics: 
\[ C_a = x_a / (m \times n), \]
\[ C_r = x_r / n, \]
\[ C_s = x_s / m \] and \( C_{ij} = x_{ij} / n \), where \(x_a\) is the number of completely specified characters\(^11\) in the MSA, \(x_r\) is
the number of completely specified characters in the \( c \)th column of the MSA and \( x_{ij} \) is the number of homologous sites with completely specified characters in both sequences (\( i \) and \( j \)). In summary, \( C_a, C_r, C_c \) and \( C_{ij} \) measure the completeness of the alignment, the \( r \)th sequence, the \( c \)th site, and the \( i \)th and \( j \)th sequences, respectively.

The first of these metrics (\( C_a \)) is related to the percent missing data used previously, but it is also, as shown in Figure 1a, the least useful completeness metric considered here: Alignments A and B differ greatly, but they have the same \( C_a \) value (i.e., 0.7). The \( C_r, C_c \) and \( C_{ij} \) metrics, on the other hand, are able to detect these differences. For example, the \( C_r \) values range from 0.3 to 1.0 for Alignment A and from 0.4 to 1.0 for Alignment B, raising greater concern, from a sequence-centric perspective, about Alignment A than about Alignment B. If we were to omit any sequence from Alignment A, then it would be sensible to omit the one with the smallest \( C_r \) value. The \( C_c \) values range from 0.2 to 1.0 for Alignment A and from 0.5 to 0.8 for Alignment B. Again, there is greater concern about Alignment A than about Alignment B (due to the lower \( C_c \) scores and the greater range of values). The \( C_{ij} \) values range from 0.3 to 1.0 for Alignment A and from 0.0 to 0.9 for Alignment B. There is cause for great concern if \( C_{ij} = 0.0 \) is detected as it means that sequences \( i \) and \( j \) have no shared homologous sites with completely specified characters in both sequences.

Evolutionary distances between such sequences cannot be estimated unless the MSA contains at least one other sequence that overlaps both \( i \) and \( j \). When such a case occurs, the evolutionary distance between sequences \( i \) and \( j \) is called inferred by proxy. Currently, the prevalence of this problem is unknown.

Figures 1b and 1c reveal the distributions of \( C_r \) and \( C_c \) for Alignments A and B, offering additional insight into the alignments’ completeness. Conveniently, the \( C_c \) scores may be used to selectively omit the least complete sites. This masking of sites in MSAs is popular in phylogenetics and many methods\(^{12-20}\) are now available. Additional information can be obtained by analyzing heat maps generated from the \( C_{ij} \) values. Figure 1d shows the heat maps obtained from Alignments A and B. The most obvious things to note are that in Alignment A Tagliatelle stands out as being the least complete sequence whereas Capellini and Spaghetti share no homologous sites with completely
specified nucleotides in both sequences in Alignment B. Although this was easy to detect in Figure 1, it will be more difficult to do if \( n \) and/or \( m \) were larger, as is typically the case in phylogenomic data.

The benefits offered by the new completeness metrics are clear, but embedding figures like those in Figure 1 in publications may be impractical. Alternatively, the essential details may be reported in a table (Table 1) or in one line (e.g., Alignment B: \( m = 10, n = 100, C_a = 0.7, C_r = [0.4, 1.0], C_c = [0.4, 0.8], \) and \( C_{ij} = [0.0, 0.9] \)). The closer to 1.0 the four \( C \) scores are, the more complete an alignment is. If, on the other hand, the values are closer to 0.0 than to 1.0, users may consider masking some of the sequences and/or sites before starting a phylogenetic analysis of the data.

Given their potential to inform researchers across a wide range of scientific disciplines, we argue that \( m, n, C_a, C_r, C_c \) and \( C_{ij} \) should be combined into what we henceforth call an MRS for MSAs, and that publications that report all of these values be labelled *compliant with the MRS for MSAs*. To our knowledge, this has never been done beforehand, leading to widespread ignorance about the MSAs that are relied upon in ground-breaking bio-medical research.

The MRS may be used to identify dubious MSAs in phylogenomic projects. These alignments occur in large phylogenomic projects because of a lack of awareness concerning a multitude of problems in the assembly, orthology assignment, and alignment procedures. Methods to identify odd MSAs are available\(^2\), but they are not yet fully reliable. The MRS enables a better outlier check for MSA.

Typically, MSAs comprise more sequences and sites than those in Figure 1a, so to facilitate using the MRS, we implemented AliStat, a fast, flexible, and user-friendly program for surveying MSAs. AliStat computes the \( C_a, C_r, C_c \), and \( C_{ij} \) values from MSAs of nucleotides, di-nucleotides, codons and amino acids., AliStat lists the results on the command-line or in files that can be accessed by other programs.
The benefit of new MRS for MSAs is underlined in two surveys of large MSAs (Table 2). In the first case, surveying an MSA of the enzyme carboxyl/cholinesterase revealed that some of the $C_r$ and $C_c$ scores are closer to 0.0 than 1.0, and that at least two sequences have no homologous sites in common with completely specified characters in both sequences. Further inspection of the output files revealed large proportions of low $C_r$ and $C_c$ scores (Supplementary material), so it might be wise to mask some of the sequences and/or sites before phylogenetic analysis of these data.

In the second case, surveying a massive concatenation of MSAs of nuclear genes revealed a more complete alignment but also low $C_r$, $C_c$, and $C_{ij}$ values. The presence of these values indicates that additional masking of this MSA might have been wise (see Supplementary material). For example, omitting the two least-complete sequences (i.e., the genera Leucoptera and Pseudopostega) could have been considered.

The MRS for MSAs is a sound solution to a large and so-far-neglected problem: how do we report, as transparently and informatively as possible, the completeness of the MSAs used in bio-medical research? Better transparency about the completeness of MSAs is clearly needed, because MSAs represent a foundational cornerstone in many bio-medical research projects and, as revealed by the example in Figure 1, MSAs may look different but have the same percentage of missing data. So far, information on the completeness of MSAs used in bio-medical research has been largely absent, leaving readers unable to critically evaluate the merits of scientific discoveries made on the basis of MSAs. It is critical to recognize, and acknowledge, that many MSAs are the result of scientific procedures. Therefore, it is necessary to present the results of these procedures more transparently and comprehensively. Many scientific papers now include links to the MSAs used, but the MSAs are often so large that it is impossible to form a comprehensive picture about the completeness of these MSAs.

Our MRS enables a radical change in scientific behavior, allowing authors to report their results more transparently, and readers the ability to critically assess discoveries made from analyses of sequence data stored in MSAs.
METHODS
Further details about AliStat are available in the online version of this paper.

DATA
Data, and code used to analyze the data, are available from http://github.com/thomaskf/AliStat.

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We thank staff at the Australian National University and University College Dublin for their helpful feedback on the color scheme used in the heat map. Many of the respondents were color-blind.

AUTHOR CONTRIBUTIONS
L.S.J. conceived the project and wrote the first version of AliStat (in C) to conduct a pilot study of the merits of using the metrics. B.M. implemented the first Perl script to produce the heat maps, and T.K.F.W. implemented the final version of AliStat (in C++). S.K., K.M., D.K. Y. and L.S.J. tested the C++ version of AliStat, and provided constructive feedback on the software. L.S.J. drafted the paper with input from the other authors.

COMPETING FINANCIAL INTERESTS
The authors declare not competing financial interests.
FIGURE LEGENDS

**Figure 1.** Example, based on two multiple sequences alignments (a), illustrating the corresponding distributions of completeness scores for rows (b), columns (c), and pairs of sequences (d).

**Table 1.** Example of the MRS for the alignments in Figure 1a

| Feature          | Alignment A | Alignment B |
|------------------|-------------|-------------|
| Sequences        | 10          | 10          |
| Sites            | 100         | 100         |
| Alphabet         | Nucleotides | Nucleotides |
| $C_a$            | 0.7         | 0.7         |
| $C_r$ [min – max] | 0.3 – 1.0  | 0.4 – 1.0  |
| $C_c$ [min – max] | 0.1 – 1.0  | 0.4 – 0.8  |
| $C_{ij}$ [min – max] | 0.3 – 1.0 | 0.0 – 0.9 |

**Table 2.** Example of the MRS for two published MSAs

| Feature          | Carboxyl/Colineesterase$^{21}$ | Lepidoptera$^{22}$ |
|------------------|-------------------------------|---------------------|
| Sequences        | 364                           | 203                 |
| Sites            | 2,645                         | 749,791             |
| Alphabet         | Amino acids                   | Amino acids         |
| $C_a$            | 0.2262                        | 0.6422              |
| $C_r$ [min – max] | 0.0106 – 0.5550               | 0.0609 – 0.9738     |
| $C_c$ [min – max] | 0.0027 – 0.9972               | 0.0000 – 0.9655     |
| $C_{ij}$ [min – max] | 0.0000 – 0.5550   | 0.0084 – 0.9672     |
The MRS for MSAs is first-of-a-kind, and AliStat, which enables compliance with the MRS for MSAs, is, to our knowledge, the first program to compute the four completeness scores presented above. AliStat is written in C++ and is available, under an CSIRO Open Source Software License Agreement (variation of the BSD / MIT License), from http://github.com/thomaskf/AliStat/.

AliStat reads a text file with nucleotide, di-nucleotide, codon or amino-acid sequences, which are aligned and saved in the FASTA format. In other words, AliStat considers alphabets with four, 16, 20, and 64 states. If the sequences comprise single nucleotides, the characters may be lumped to form six 3-state alphabets (i.e., CRT, AGY, ACK, GMT, AST and CGW) and seven 2-state alphabets (i.e., RY, KM, SW, AB, CD, GH and TV). If the 3- and 2-state alphabets are used, the letters R, Y, K, M, S, W, B, D, H and V are considered completely specified characters (unlike normal practice\textsuperscript{11}).

AliStat can be run in two modes: Brief mode or Full mode. Execution in brief mode is done using the following command:

\texttt{alistat <infile> <data type> -b}

and results in the following output format are printed to the terminal:

File name, #seqs, #sites, $C_a$, max $C_r$, min $C_r$, max $C_c$, min $C_c$, max $C_{ij}$, min $C_{ij}$

The brief-mode execution was included to allow users to quickly obtain the essential values from a great number of alignments (e.g., when comparing genomes phylogenetically).

The full-mode execution (default option) allows other options to be used and is intended when a more detailed examination of an MSA is required. For example, the $-t$ option is used to indicate what types of $C$ scores should be printed in output files, the $-m$ option is used to set a threshold for masking sites, and the $-i$ option is used to indicate that a heat map is needed. Other options
and how all of the options may be used are described in the AliStat manual. The same information can be obtained by typing

```
alistat -h
```

in the command-line.

The output files appear in the .txt, .csv, .R, .dis, .svg, and .fst formats, which can be processed by other software packages. The .txt file summarizes the results. The .csv files present the $C$ scores and may be examined using R. For example, if a user wishes to generate a histogram of the $C_c$ scores, the Table_2.csv file may be analyzed using the Histogram_Cr.R file. In some cases, users may want to infer a tree or network based on the $C_{ij}$ score (or the $I_{ij}$ score, where $I_{ij} = 1.0 - C_{ij}$). In such cases, .dis files may be analyzed by, for example, SplitsTree\textsuperscript{23}. The heat map, which may be triangular or square, is stored in the .svg file and may be opened using Adobe Illustrator\textsuperscript{\textregistered}. If the $-m$ option is used, the original MSA is split into two, with all sites having a $C_c$ score larger than a user-specified threshold saved in a file called Mask.fst and the other sites saved in a file called Disc.fst. The two .fst files may be analyzed separately by other means (e.g., phylogenetic programs).
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**Alignment A**

| Linguine    | TCGCAGGATCGTATAGGAGGTGTCTTACGGCCAATATAAGAGACGCCGTAAGAGTGATTCTGCAAGCCAACCTAAGCCGTTTAAGAGCTGGGCGTTTACCGCTGCTGCTC |
|-------------|----------------------------------------------------------------------------------------------------------------|
| Tagliatelle | --------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Fettuccine  | ---------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Capellini   | ---------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Spaghetti   | ---------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Vermicelli  | ---------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Lasagne     | ---------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Farfallini  | ---------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Tortellini  | ---------------------------------------------------------------------------------------------------------------------------------------------------------------------- |
| Ravioli     | GCGTGGAGCGGCCCCTGAAAACGACCACCCACCAAGTTTTCTGAAAGAGATGCTGATGGCGCTTC |  

**Alignment B**

| Linguine    | CCAATGAACAAAACCGACCCAGGCCGACACGGTGA                                                                                                                                     |
|-------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Tagliatelle | GCAGGGCTTTTTTAGCCAGCATA                                                                                                                                                    |
| Fettuccine  | GCAGGGCGGCTTTTTAGCCAGCCATA                                                                                                                                                    |
| Capellini   | CAGCACGTGCTGACATGGTTTTCTGAAAGAGATGCTGATGGCGCTTC                                                                                                                             |
| Spaghetti   | CACACGTGCTGACATGGTTTTCTGAAAGAGATGCTGATGGCGCTTC                                                                                                                             |
| Lasagne     | CCAACGTGCTGACATGGTTTTCTGAAAGAGATGCTGATGGCGCTTC                                                                                                                             |
| Farfallini  | CACACGTGCTGACATGGTTTTCTGAAAGAGATGCTGATGGCGCTTC                                                                                                                             |
| Tortellini  | CACACGTGCTGACATGGTTTTCTGAAAGAGATGCTGATGGCGCTTC                                                                                                                             |
| Ravioli     | CACACGTGCTGACATGGTTTTCTGAAAGAGATGCTGATGGCGCTTC                                                                                                                             |

**Frequency**

Alignment A

- Cr: 0.0 1.0
- Cc: 0.0 1.0

Alignment B

- Cr: 0.0 1.0
- Cc: 0.0 1.0

**Heatmaps**

- Alignment A
- Alignment B

**Legend**

- ≤ 1.0
- < 0.7
- < 0.5
- < 0.3
- < 0.1
- = 0.0
Supplementary Material:

A minimum reporting standard for multiple sequence alignments

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Analysis of an alignment of carboxyl/cholinesterases (CCEs) from a paper by Pearce et al.\textsuperscript{1}

The alignment of amino acids used to annotate the CCE genes from Helicoverpa armigera, H. zea, Manduca sexta and Bombyx mori was surveyed using AliStat v1.11. The alignment comprised 364 sequences and 2645 sites. **Figure S1** presents the distribution of $C_r$ values, with several sequences having values close to 0.0. Because the objective of the study by Pearce et al.\textsuperscript{1} was to annotate the genes, it was not possible to remove any sequences from the data.

**Figure S1.** Histogram showing the distribution of $C_r$ scores from the CCE genes. The arrows point to the lowest $C_r$ scores.

**Figure S2** reveals the distribution of $C_c$ scores, with a high proportion of sites with low $C_c$ values. Based on this distribution, sites with $C_c \leq 0.5$ were masked in the study by Pearce et al.\textsuperscript{1}.
Figure S2. Histogram showing the distribution of $C_c$ scores from the CCE genes.

Figure S3 shows the heat map of $C_{ij}$ scores, revealing a high proportion of sequence pairs with low $C_{ij}$ values. Because the aim of the study by Pearce et al.\textsuperscript{1} was to annotate the genes, it was not possible to remove any sequences from the data.

Figure S3. Heat map showing the distribution of $C_{ij}$ scores from the CCE genes. The benefit of this heat map is realized by enlarging the image, at which point it become obvious that two sequences, labelled HzeaCCE016h and MsexCCE001s, have no homologous sites with unambiguous characters in both sequences.

Pearce et al.\textsuperscript{1} used a threshold of $C_c = 0.5$ to mask the alignment of amino acids. Table 1 reveals the effect of doing so. In this case, the table complies with the minimum reporting standard (MRS) for multiple sequence alignments (MSAs).
Table 1. Example highlighting the effect of using $C_c = 0.5$ to mask the sites in the alignment of CCEs.

| Feature      | Before masking | After masking |
|--------------|----------------|--------------|
| Sequences    | 364            | 364          |
| Sites        | 2,645          | 546          |
| Alphabet     | Amino acids    | Amino acids  |
| $C_a$        | 0.2262         | 0.9562       |
| $C_r$ [min – max] | 0.0106 – 0.5550 | 0.0458 – 0.9890 |
| $C_c$ [min – max] | 0.0027 – 0.9972 | 0.5055 – 0.9972 |
| $C_{ij}$ [min – max] | 0.0000 – 0.5550 | 0.0000 – 0.9890 |

Analysis of lepidopteran nuclear data from a study by Kawahara et al.²

The alignment of amino acids labelled SA4_aminoacid_supermatrix_resorted_renamed.fas was surveyed using AliStat v1.11. The alignment comprised 203 sequences and 749,791 sites. Figure S4 reveals the distribution of $C_r$ values, with most sequences having values close to 1.0. In this case, only a few sequences had very low $C_r$ scores.

Figure S4. Histogram showing the distribution of $C_r$ scores from the super-alignment (i.e., 2098 MSAs of concatenated single-copy genes). The arrows point to the two lowest $C_r$ scores.

Figure S5 reveals the distribution of $C_c$ scores, with a high proportion of sites with high $C_c$ values. Based on this distribution, omitting sites with $C_c \leq 0.2$ might have been sufficient.
Figure S5. Histogram showing the distribution of $C_c$ scores from the super-alignment (Kawahara et al.2).

Figure S6 shows the heat map of $C_{ij}$ scores, revealing a low proportion of sequence pairs with low $C_{ij}$ values. In this case, the sequences found to be least complete are from the genera *Leucoptera* and *Pseudopostega*.
Figure S6. Heat map showing the distribution of $C_{ij}$ scores from the concatenate gene alignments. Again, the benefit of this heat map is realized by enlarging the image. In this case, the two most incomplete sequences are identified as being from the genera *Leucoptera* and *Pseudopostega*.

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