NOMENCLATURE RECOMMENDATIONS

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Nomenclature for the description of human sequence variations

Received: 12 March 2001 / Accepted: 13 March 2001 / Published online: 19 June 2001 © Springer-Verlag 2001

Abstract A nomenclature system has recently been suggested for the description of changes (mutations and polymorphisms) in DNA and protein sequences. These nomenclature recommendations have now been largely accepted. However, current rules do not yet cover all types of mutations, nor do they cover more complex mutations. This document lists the existing recommendations and summarizes suggestions for the description of additional, more complex changes. Another version of this paper has been published in Hum Mut 15:7–12, 2000.

Introduction

Recently, a nomenclature system has been suggested for the description of changes (mutations and polymorphisms) in DNA and protein sequences (Antonarakis and the Nomenclature Working Group 1998, http://www.dmd. nl/mutnomen.html). These nomenclature recommendations have now been largely accepted and stimulated the uniform and unequivocal description of sequence changes. However, current rules do not yet cover all types of mutations, nor do they cover more complex mutations. This document lists the existing recommendations and summarizes suggestions for the description of additional and more complex changes (shown in italics) based on den Dunnen and Antonarakis (2000).

Discussions regarding the advantages and disadvantages of the suggestions are necessary in order to continuously improve the designation of sequence changes. The consensus of the discussions will be posted on a WWWpage (http://www.dmd.nl/mutnomen.html), and we invite

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Division de Génétique Médicale, Centre Médical Universitaire, Rue Michel-Servet 1, 1211 Genève, Switzerland e-mail: stylianos.antonarakis@medecine.unige.ch, Fax: +41-22-702-5706 investigators to communicate with us regarding these suggestions. Furthermore, we invite investigators to send us complicated cases not covered yet, with a suggestion of how to describe these (mail to ddunnen@lumc.nl and Stylianos.Antonarakis@medecine.unige.ch). We hope the WWW-pages will be used as a guide to describe any sequence change, ultimately evolving into a uniformly accepted standard.

General recommendations

Suggestions extending the current recommendations are in italtics.

The term "*sequence variation*" is used to prevent confusion with the terms "*mutation*" and "*polymorphism*", mutation meaning "change" in some disciplines and "disease-causing change" in others and polymorphism meaning "non-disease-causing change" or "change found at a frequency of 1% or higher in the population".

The basic recommendation is to use *systematic names* to describe each sequence variation. For this, variations are described at the most basic level, i.e., the DNA level, using either a genomic or a cDNA reference sequence. A genomic reference sequence is preferred because it overcomes difficult cases, including multiple transcription initiation sites (promoters), alternative splicing, the use of different poly-A addition signals, multiple translation initiation sites (ATG-codons), and the occurrence of length variations. When, like in most cases, the entire genomic sequence is not known, a cDNA reference sequence should be used instead.

- Sequence variations are described in relation to a reference sequence for which the accession number from a primary sequence database (Genbank, EMBL, DDJB, SWISS-PROT) should be mentioned in the publication/database submission (e.g., M18533)
- Tabular listings of the sequence variations described should contain columns for DNA, RNA, and protein and clearly indicate whether the changes were **experimentally determined** or only theoretically deduced

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- To avoid confusion in the description of a sequence change, preceed the description with a letter indicating the type of reference sequence used:
 - "g." for a *genomic* sequence (e.g., g.76A>T)
 - "c." for a *cDNA* sequence (e.g., c.76A>T)
 - "m." for a *mitochondrial* sequence (e.g., m.76A>T)
 - "r." for an **RNA** sequence (e.g., r.76a>u)
 - "p." for a protein sequence (e.g., p.K76A)
- To discriminate between the different levels (DNA, RNA, or protein), descriptions are unique:
 - at DNA-level, in capitals, starting with a number referring to the first nucleotide affected (e.g., c.76A>T)
 - at RNA-level, in lower-case, starting with a number refering to the first nucleotide affected (e.g., r.76a>u)
 - at protein level, in capitals, starting with a letter referring to the first amino acid (one-letter code) affected (e.g., p.T26P)
- A range of affected residues is indicated by a "_"character (underscore) separating the first and last residue affected (e.g., 76_78delACT)

NOTE: current recommendations use the "-"-character (i.e., 76–78delACT)

- For deletions or duplications in single nucleotide (or amino acid) stretches or tandem repeats, the most 3' copy is arbitrarily assigned to have been changed (i.e. ACTTTG<u>TG</u>CC to ACTTTGCC is described as 7_8delTG)
- Two sequence variations in one allele are listed between brackets, *separated by a ";"-character (e.g.,* [76A>C; 83G>C])

NOTE: the recommendations made in den Dunnen and Antonarakis (2000) to use a "+"-character as a separator (i.e., [76A>C+ 83G>C]) has been retracted

- Sequence changes in different alleles (e.g., for recessive diseases) are listed between brackets, separated by a "+"-character (e.g., [76A>C] + [87delG]) NOTE: the current recommendation is [76A>C + 87delG]
- A unique identifier should be assigned to each mutation. The unique OMIM-identifier can be used, otherwise database curators should assign unique identifiers

DNA level

- Nucleotides are designated by the bases (in upper case); A (adenine), C (cytosine), G (guanine), and T (thymidine)
- Nucleotide numbering:
 - nucleotide +1 is the A of the ATG-translation initiation codon, the nucleotide 5' to +1 is numbered -1; there is no base 0

 \circ non-coding regions:

the nucleotide 5' of the ATG-translation initiation codon is -1

the nucleotide 3' of the translation termination codon is *1

 \circ intronic nucleotides:

beginning of the intron: the number of the last nucleotide of the preceeding exon, a plus sign, and the position in the intron, e.g., 77+1G, 77+2T (when the exon number is known, the notation can also be described as IVS1+1G, IVS1+2T)

end of the intron: the number of the first nucleotide of the following exon, a minus sign, and the position upstream in the intron, e.g., 78–2A, 78–1G (when the exon number is known, the notation can also be described as IVS1–2A, IVS1–2G)

Description of nucleotide changes

- Substitutions are designated by a ">"-character
 - 76A>C denotes that at nucleotide 76 an A is changed to a C
 - 88+1G>T (alternatively IVS2+1G>T) denotes the G to T substitution at nucleotide +1 of intron 2, relative to the cDNA positioned between nucleotides 88 and 89
 - 89–2A>C (alternatively IVS2–2A>C) denotes the A to C substitution at nucleotide –2 of intron 2, relative to the cDNA positioned between nucleotides 88 and 89

NOTE: *polymorphic variants* are sometimes described as 76A/G, but this is not recommened!

- **Deletions** are designated by "del" after the nucleotide(s) flanking the deletion site
- 76_78del (alternatively 76_78delACT) denotes a ACT deletion from nucleotides 76 to 78
- 82_83del (alternatively 82_83delTG) denotes a TG deletion in the sequence ACTTTG<u>TG</u>CC (A is nucleotide 76) to ACTTTGCC
- IVS2_IVS5del (alternatives 88+?-923-? or EX3_5del) denotes an exonic deletion starting at an unknown position in intron 2 (after cDNA nucleotide 88) and ending at an unknown position in intron 5 (before cDNA nucleotide 923)
- **Duplications** are designated by "dup" after the first and last nucleotide affected by the duplication
 - 77–79dup (or 77_79dupCTG) denotes that the nucleotides 77 to 79 were duplicated
 - duplicating insertions in single nucleotide stretches (or short tandem repeats) are preferably described as a duplication, e.g., a TG insertion in the TG-tandem repeat sequence of ACTTTGTGCC (A is nt 76) to ACTTTGTG<u>TGCC</u> is described as 82_83dupTG (now 83_84insTG)

• <u>Insertions</u> are designated by "ins" after the nucleotides flanking the insertion site, followed by the nucleotides inserted

NOTE: as separator the "^"-character is sometimes used but this is not recommened (e.g., 83^84insTG)

- 76_77insT denotes that a T was inserted between nucleotides 76 and 77
- 83_84dupTG denotes a TG insertion in the TG-tandem repeat sequence of ACTTTGTGCC (A is nucleotide 76) to ACTTTGTG<u>TG</u>CC (*see "duplications"*)
- Variability of short sequence repeats, e.g., in ACT-GTGTGCC (A is nt 1991), are designated as 1993(TG)3–6 with nucleotide 1993 containing the first TG-dinucleotide, which is found repeated 3 to 6 times in the population.
- <u>Insertion/deletions (indels)</u> are described as a deletion followed by an insertion after the nucleotides affected
 112_117delinsTG (alternatively 112_117delAGGT-CAinsTG or 112_117>TG) denotes the replacement of nucleotides 112 to 117 (AGGTCA) by TG
- <u>Inversions</u> are designated by "inv" after the first and last nucleotides affected by the inversion
 - 203_506inv (or 203_506inv304) denotes that the 304 nucleotides from position 203 to 506 have been inverted
- <u>Translocations</u> (no suggestions yet)
- <u>Changes in different alleles</u> (e.g., in recessive diseases) are described as "[change allele 1] + [change allele 2]"
 - [76A>C] + [76A>C] denotes a homozygous A to C change at nucleotide 76
 - \circ [76A>C] + [?] denotes an A to C change at nucleotide 76 in one allele and an unknown change in the other allele
- <u>*Two variations in one allele</u>* are described as "[first change + second change]"</u>
 - [76A>C; 83G>C] denotes an A to C change at nucleotide 76 and a G to C change at nucleotide 83 in the same allele

RNA level

Sequence changes at the RNA level are basically described as those at the DNA level with the following modifications/additions:

- An "r." is used to indicate that a change is described at RNA-level
- *Nucleotides are designated by the bases (in lower case); a (adenine), c (cytosine), g (guanine), and u (uracil)*
 - 78*u*>*a* denotes that at nucleotide 78 *a U* is changed to an A

- When one change affects RNA-processing, yielding two or more transcripts, these are described between square brackets, separated by a ","-character
 - [r.76a>c, r.76a>c; r.73_88del] denotes the nucleotide change c.76A>C causing the appearance of two RNA molecules, one carrying this variation only and one containing in addition a deletion of nucleotides 73 to 88 (shift of the splice donor site to within the exon)
 - \circ [*r*=, *r*.88_89ins88+1_88+10; *r*.88+2t>c] denotes the intronic mutation *c*.88+2T>C causing the appearance of two RNA molecules, one normal (*r*=) and one containing an insertion of the intronic nucleotides 88+1 to 88+10 with the nucleotide change 88+2t>c
 - [r.88g>a; r.88_89ins88+1_88+10] denotes the nucleotide change c.88G>A causing an insertion of the intronic nucleotides 88+1 to 88+10 (shift of the splice donor site to an intronic position)

Protein level

Sequence changes at protein level are basically described as those at the DNA level with the following modifications/additions:

- The one letter amino acid code is used, with "X" designating a translation termination codon
- Amino acid numbering;

the translation initiator Methionine is numbered as +1

Description of amino acid changes

• Substitutions;

- *missense changes* W26C denotes that amino acid 26 (Tryptophan, W) is changed to a Cysteine (C) NOTE: polymorphic variants are sometimes described as 36L/I, but this is not recommend!
- *nonsense changes* W26X denotes that amino acid
 26 (Tryptophan, W) is changed to a stop codon (X)
- \circ initiating methionine (M1)

Currently, mutations in the translation initiating Methionine (M1) are mostly described as a substitution, e.g., M1 V. This is not correct. Either no protein is produced or the translation initiation site moves upor downstream. Unless experimental proof is available, it is probably best to report the effect on protein level as "p.?" (unknown). When experimental data show that no protein is made, the description "p.0" might be most appropriate

- **Deletions** are designated by "del" after the nucleotide(s) flanking the deletion site
 - K29del in the sequence C<u>K</u>MGHQQQCC (C is amino acid 28) denotes a deletion of amino acid Lysine 29 (K) to CMGHQQQCC
 - C28_M30del denotes a deletion of three amino acids, from Cysteine 28 to Methionine 30

- Q35del in the sequence CKMGHQQQCC (C is amino acid 28) denotes a Glutamine 35 (Q) deletion to CKMGHQQCC
- if a deletion creates a new amino acid at the deletion junction the change is described as an <u>inser-</u> <u>tion/deletion</u>, e.g., C28_M30delinsW (see below)
- **Duplications** are designated by "dup" after the first and last amino acid affected by the duplication
 - G31_Q33dup in the sequence CKMGHQQQCC (C is amino acid 28) denotes a duplication of amino acids Glycine 31 (G) to Glutamine 33 (Q) CK-MGHQGHQQQCC
 - duplicating insertions in single amino acid stretches (or short tandem repeats) are described as a duplication, e.g., an HQ insertion in the HQ-tandem repeat sequence of CKMGHQHQCC (C is amino acid 28) to CKMGHQHQHQCC is H34_Q35dup (now Q35_C36insHQ)
- <u>Insertions</u> are designated by "ins" after the nucleotides flanking the insertion site, followed by the nucleotides inserted

NOTE: as separator the "^"-character is sometimes used but this is not recommened (e.g., Q83^C84insQ)

- K29_M29insQSK denotes that the sequence QSK was inserted between amino acids Lysine 29 (K) and Methionine 30 (M), changing CKMGHQQQCC (C is amino acid 28) to CK<u>OSK</u>MGHQQQCC
- Q35dup in the sequence CKMGHQQQCC (C is amino acid 28) denotes a duplicating insertion of a Glutamine (Q) to CKMGHQQQQCC (see "duplications")
- if an insertion creates a new amino acid at the insertion junction the change is described as an <u>in-</u> <u>sertion/deletion</u>, e.g., C28delinsWV (see below)

- Variability of short sequence repeats, e.g., in CK-MGHQQQCC (C is amino acid 28), are designated as 33(Q)3–6 with amino acid Glutamine 33 (Q, the first repeated amino acid) found repeated 3 to 6 times in the population.
- <u>Insertion/deletions (indels)</u> are described as a deletion followed by an insertion after the nucleotides affected
 - C28_K29delinsW denotes a 3 bp deletion affecting the codons for Cysteine 28 and Lysine 29, substituting them for a codon for Tryptophan
 - C28delinsWV denotes a 3 bp insertion in the codon for Cysteine 28, generating codons for Tryptophan (W) and Valine (V)
- <u>Frame shifting mutations</u>; recommendations to describe these sequence changes have not yet been made. Although it is probably not useful to add much detail in this description, it might be sensible, e.g., in the case of C-terminal mutations, to include the length of the new, shifted reading frame
 - R97fsX121 (alternative R97Xfs) denotes a frame shifting change with Arginine 97 as the first affected amino acid and the new reading frame being open for 23 amino acids

References

- Antonarakis SE, Nomenclature Working Group (1998) Recommendations for a nomenclature system for human gene mutations. Hum Mutat 11:1–3
- Dunnen JT den, Antonarakis SE (2000) Mutation nomenclature extensions and suggestions to describe complex mutations: a discussion. Hum Mutat 15:7–12 (copy in PDF-format) http://www.dmd.nl