GENETIC NOMENCLATURE

Strain collections. The ease of rapidly accumulating a large number of mutants requires careful bookkeeping to avoid confusing one mutant with another. Each mutant should be assigned a strain number. Strain numbers usually consist of 2-3 capital letters designating their source and a serial numbering of the strains in a central laboratory collection. It is a good idea to check existing genetic resources to avoid the potential confusion that can result from assigning different genes the same name. In addition to genome databases, good resources for gene names include the Salmonella Genetic Stock Centre (http://www.ucalgary.ca/~kesander/), and the *E. coli* Genetic Stock Center (http://cgsc.biology.yale.edu/).

Nomenclature. Through the 1960’s, genetic nomenclature was a “tower of babel”. Due to the absence of clear rules for naming genes, each investigator assigned new names haphazardly, often resulting in the same name being applied to different genes or different names being applied to the same gene. To further confuse the issues, different investigators would each assign allele numbers independently, so two different alleles might have the exact same designation. To eliminate the resulting confusion, Demerec et al. (1966, 1986) developed a standard nomenclature for bacterial genes. With the development of new genetic tools, some modifications have been required. A detailed description of these rules can be found in the instructions to authors for the *J. Bacteriol.* (http://jb.asm.org/misc/ifora.shtml). The basic rules are described below.

GENOTYPE:

1. Genes. Each gene is assigned a three-letter designation, usually an abbreviation for the pathway or the phenotype of mutants. When the genotype is indicated, the three-letter designation is written in lower case. Different genes that affect the same pathway are distinguished by a capital letter following the three-letter designation.

   For example, mutations affecting pyrimidine biosynthesis are designated *pyr*; the *pyrC* gene encodes the enzyme dihydroorotase and the *pyrD* gene encodes the enzyme dihydroorotate dehydrogenase.

2. Allele numbers. Each mutation in the pathway is consecutively assigned a unique allele number. A separate series of allele numbers is used for each three-letter locus designation. If there is no capital letter designating a specific gene, insert a dash before the allele number. Blocks of allele numbers are assigned to laboratories by the appropriate genetic stock center.

   For example, *pyrC19* refers to a particular *pyr* mutation that affects the *pyrC* gene. In order to distinguish each mutation, no other *pyr* mutation, regardless of the gene affected, will be assigned the allele number 19. A separate series of allele numbers is used for each three-letter locus designation. Allele numbers should be used sequentially and carefully monitored to insure that two different mutations are not named with the same allele numbers.

   The entire genotype is italicized or underlined (e.g. *pyrC19*).

3. Insertions. Transposable elements or suicide plasmids can insert in known genes or in a site on the chromosome where no gene is yet known. When an insertion is in a known gene, the mutation is given a three-letter designation, gene designation, and allele number as described above, followed by a double colon then the type of insertion element. DO NOT leave blank spaces between the letters or numbers and the colon. For example, a particular Tn10 insertion within the *pyrC* gene (mutant allele number 103) may be designated *pyrC103::Tn10*. 

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When a transposon insertion is not in a known gene, it is named according to the map position of the insertion on the chromosome. Such insertions are named with a three-letter symbol starting with z. The second and third letters indicate the approximate map position in minutes: the second letter corresponds to 10-minute intervals of the genetic map numbered clockwise from minute 0 (a = 0-9; b = 10-19; c = 20-29, etc.); the third letter corresponds to minutes within any 10-minute segment (a = 0; b = 1; c = 2; etc.). For example, a Tn10 insertion located near pyrC at 23 minutes is designated zcd::Tn10. Allele numbers are assigned sequentially to such insertions regardless of the letters appearing in the second and third positions, so if more refined mapping data suggests a new three-letter symbol, the allele number of the insertion mutation is retained. This nomenclature uses zaa (0 min) to zjj (99 min). Insertion mutations on extrachromosomal elements are designated with zz, followed by a letter denoting the element used. For example, zzf is used for insertion mutations on an F\textsuperscript{\textprime} plasmid. Insertions with an unknown location are designated zxx.

\begin{align*}
zaa &= \text{insertion at 0-1 min} \\
zab &= \text{insertion at 1-2 min} \\
zac &= \text{insertion at 2-3 min} \\
zad &= \text{insertion at 3-4 min} \\
zae &= \text{insertion at 4-5 min} \\
zaf &= \text{insertion at 5-6 min} \\
zag &= \text{insertion at 6-7 min} \\
zah &= \text{insertion at 7-8 min} \\
zai &= \text{insertion at 8-9 min} \\
zaj &= \text{insertion at 9-10 min} \\
zxx &= \text{insertion with unknown location} \\
zzf &= \text{insertion on F-plasmid}
\end{align*}

Some commonly used mini-transposon derivatives are designated as follows:

\begin{align*}
\text{Tn10dTet} &= \text{Tet resistance, deleted for Tn10 transposase} \\
\text{Tn10dCam} &= \text{Derived from Tn10dTet, Cam resistance substituted for Tet resistance} \\
\text{Tn10dKan} &= \text{Derived from Tn10dTet, Kan resistance substituted for Tet resistance} \\
\text{Tn10dGen} &= \text{Derived from Tn10dTet, Gen resistance substituted for Tet resistance} \\
\text{MudJ} &= \text{Kan resistance, forms } \textit{lac} \text{ operon fusions, deleted for Mu transposase} \\
\text{MudJ-Cam} &= \text{Derived from MudJ, Cam resistance marker disrupts Kan resistance} \\
\text{MudCam} &= \text{Cam resistance substitution between ends of Mu}
\end{align*}

4. **Plasmids.** Plasmids should be indicated by a / slash after the genotype. Indicate the name of the plasmid, the plasmid origin, and the relevant genotype or phenotype carried by the plasmid.

Insertions of suicide plasmids into the chromosome can be indicated as described for transposons. If a duplication is generated it can be described as indicated under chromosomal rearrangements.

5. **Phage.** Prophages or plasmids integrated into an attachment site can be indicated by the name of the attachment site followed by a double colon and the phage genotype indicated in brackets. For example, \text{att::[P22 mnt::Kan]}. 

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6. **Chromosome rearrangements.** Chromosome rearrangements including deletions, duplications, and inversions should be indicated by a three letter symbol indicating the type of rearrangement, followed by the genes involved indicated in parenthesis, followed by the allele number.

- **Deletions** = DEL(genes)allele number
- **Inversions** = INV(join point gene #1 – join point gene #2)allele number
- **Duplications** = DUP(gene #1*join point*gene #2)allele number

**PHENOTYPE:**

1. **Growth phenotypes.** It is often necessary to distinguish the phenotype of a strain from its genotype. The phenotype is usually indicated with the same three-letter designation as the genotype but phenotypes start with capital letters and are not underlined. (For example, strain TR251 [hisC527 cysA1349 supD] has a Cys+ His+ phenotype because the supD mutation suppresses the amber mutations in both the cysA and the hisC genes.)

2. **Antibiotic resistance.** Both two and three letter designations are commonly used for antibiotic resistance markers. Both are acceptable, but it is essential to be consistent. Resistance and sensitivity is indicated with a superscript but on the computer it is often simpler to indicate resistance with (R) and sensitivity with (S).

   - **Amp** = Ampicillin
   - **Cam** = Chloramphenicol
   - **Gen** = Gentamicin
   - **Kan** = Kanamycin
   - **Neo** = Neomycin
   - **Spc** = Spectinomycin
   - **Str** = Streptomycin
   - **Tet** = Tetracycline
   - **Zeo** = Zeomycin
   - **XG** = X-gal
   - **XP** = X-phosphate

3. **Conditional alleles.** Conditional alleles indicated by the genotype including allele number followed by the two letter designation for the conditional phenotypes shown in parenthesis. For example, *leuA414*(Am). Note that because this is a phenotype it begins with a capital letter.

   - (Ts) = Temperature sensitive mutation
   - (Cs) = Cold sensitive mutation
   - (Am) = Amber mutation
   - (Op) = Opal mutation
   - (Oc) = Ochre mutation

**REFERENCES:**
