## **SLiM Reference Sheet**

```
slim -version | -usage | -testEidos | -testSLiM |
  [-seed <seed>] [-time] [-mem] [-Memhist] [-long [<l>]] [-x] [-define <def>] <script file>
```

Types: N:NULL, l:logical, i:integer, f:float, n:numeric, s:string, o<X>:object of class X

```
(ret-type)functionName(params) { ... }
                                                                               user-defined function
[<id>] initialize() { ... }
                                                                               initialize() callback
[<id>] [gen1 [: gen2]] [early()] { ... }
                                                                               early() Eidos event
[<id>] [gen1 [: gen2]] late() { ... }
                                                                               late() Eidos event
[<id>] [gen1 [: gen2]] fitness(<mutTypeId> [, <subpopId>]) { ... }
                                                                               fitness() callback
[<id>] [gen1 [: gen2]] interaction(<intTypeId> [, <subpopId>]) { ... }
                                                                               interaction() callback
[<id>] [gen1 [: gen2]] mateChoice([<subpopId>]) { ... } (WF)
                                                                               mateChoice() callback
[<id>] [gen1 [: gen2]] modifyChild([<subpopId>]) { ... }
                                                                               modifyChild() callback
[<id>] [gen1 [: gen2]] recombination([<subpopId>]) { ... }
                                                                               recombination() callback
[<id>] [gen1 [: gen2]] mutation([<mutTypeId> [, <subpopId>]]) { ... }
                                                                               mutation() callback
[<id>] [gen1 [: gen2]] reproduction([<subpopId> [, <sex>]]) ... (nonWF)
                                                                               reproduction() callback
```

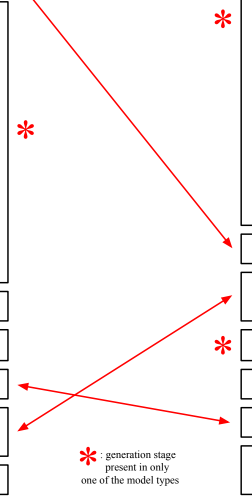
The sequence of events within one generation in WF models.

The sequence of events within one generation in nonWF models.

- 1. Execution of early() events
  - 2. Generation of offspring:
- 2.1. Choose source subpop
- 2.2. Choose parent 1
- 2.3. Choose parent 2 (mateChoice() callbacks)
- 2.4. Generate the offspring (including mutation() and recombination() callbacks)
- 2.5. Suppress/modify child (modifyChild() callbacks)
- 3. Removal of fixed mutations
- 4. Offspring become parents
- 5. Execution of late() events
- 6. Fitness value recalculation using fitness() callbacks
- 7. Generation count increment

1. Generation of offspring:

- 1.1. Call reproduction() callbacks for individuals
- 1.2. The callback(s) make calls requesting offspring
- 1.3. Generate the offspring (including mutation() and recombination() callbacks)
- 1.4. Suppress/modify child (modifyChild() callbacks)
- 2. Execution of early() events
- 3. Fitness value recalculation using fitness() callbacks
- 4. Viability/survival selection
- 5. Removal of fixed mutations
- 6. Execution of late() events
- 7. Generation count increment, individual age increments



```
Initialization functions (callable only from initialize() callbacks):
(integer$)initializeAncestralNucleotides(is seguence) (nuc)
(void)initializeGeneConversion(n$ nonCrossoverFraction, n$ meanLength,
      n$ simpleConversionFraction, [n$ bias])
(o<GElement>)initializeGenomicElement(io<GEType> genomicElementType, i start, i end)
(o<GEType>$)initializeGenomicElementType(is$ id, io<MutType> mutationTypes, n proportions,
      [Nf mutationMatrix])
(void)initializeHotspotMap(numeric multipliers, [Ni ends], [s$ sex]) (nuc)
(o<IntType>$)initializeInteractionType(is$ id, s$ spatiality, [l$ reciprocal],
      [n$ maxDistance], [s$ sexSegregation])
(void)initializeMutationRate(n rates, [Ni ends], [s$ sex])
(o<MutType>$)initializeMutationType(is$ id, n$ dominanceCoeff, s$ distributionType, ...)
(o<MutType>$)initializeMutationTypeNuc(is$ id, n$ dominanceCoeff, s$ distributType, ...) (nuc)
(void)initializeRecombinationRate(n rates, [Ni ends], [s$ sex])
(void)initializeSex(s$ chromosomeType, [n$ xDominanceCoeff])
(void)initializeSLiMModelType(s$ modelType)
(void)initializeSLiMOptions([l$ keepPedigrees], [s$ dimensionality], [s$ periodicity],
      [i$ mutationRuns], [l$ preventIncidentalSelfing], [l$ nucleotideBased])
(void)initializeTreeSeq([l$ recordMutations], [Nif$ simplificationRatio],
      [Ni$ simplificationInterval], [l$ checkCoalescence], [l$ runCrosschecks])
fitness():
                                modifyChild():
                                                                      mateChoice(): (WF)
                                                                        individual (o<Ind>$)
  mut (o<Mutation>$)
                                  child (o<Ind>$)
                                  childGenome1 (o<Genome>$)
  homozygous (l$)
                                                                        genome1 (o<Genome>$)
  relFitness (f$)
                                  childGenome2 (o<Genome>$)
                                                                        genome2 (o<Genome>$)
  individual (o<Ind>$)
                                                                        subpop (o<Subpop>$)
                                  childIsFemale (l$)
  genome1 (o<Genome>$)
                                  parent1 (o<Ind>$)
                                                                        sourceSubpop (o<Subpop>$)
  genome2 (o<Genome>$)
                                  parent1Genome1 (o<Genome>$)
                                                                        weights (f)
                                  parent1Genome2 (o<Genome>$)
  subpop (o<Subpop>$)
                                  isCloning (l$)
                                                                      recombination():
reproduction(): (nonWF)
                                  isSelfing (l$)
                                                                        individual (o<Ind>$)
                                  parent2 (o<Ind>$)
  individual (o<Ind>$)
                                                                        genome1 (o<Genome>$)
                                  parent2Genome1 (o<Genome>$)
  genome1 (o<Genome>$)
                                                                        genome2 (o<Genome>$)
                                  parent2Genome2 (o<Genome>$)
  genome2 (o<Genome>$)
                                                                        subpop (o<Subpop>$)
                                  subpop (o<Subpop>$)
  subpop (o<Subpop>$)
                                                                        breakpoints (i)
                                  sourceSubpop (o<Subpop>$)
                                                                        gcStarts (i)
Individual (Ind):
                                                                        gcEnds (i)
age <-> (i$) (nonWF)
color <-> (s$)
                                                                      mutation():
fitnessScaling <-> (f$)
                                                                        mut (o<Mutation>$)
genomes => (o<Genome>)
                                                                        genome (o<Genome>$)
genome1 => (o<Genome>$)
                                                                        element (o<GElement>$)
genome2 => (o<Genome>$)
                                                                        originalNuc (i$)
index => (i\$)
                                                                        parent (o<Ind>$)
migrant => (l$)
                                                                        subpop (o<Subpop>$)
pedigreeID => (i$)
pedigreeParentIDs => (i)
                                                                      interaction():
pedigreeGrandparentIDs => (i)
                                                                        distance (f$)
sex => (s$)
                                                                        strength (f$)
spatialPosition => (f)
                                                                        exerter (o<Ind>$)
subpopulation => (o<Subpop>$)
                                                                        receiver (o<Ind>$)
tag \leftarrow (i$)
                                                                        subpop (o<Subpop>$)
tagF <-> (f$)
uniqueMutations => (o<Mut>)
x \leftarrow > (f\$)
\vee \leftarrow > (f\$)
z \leftarrow > (f$)
                                                                      SLiM globals:
- (l)containsMutations(o<Mut> mutations)
                                                                         sim (o<SLiMSim>$)
- (i$)countOfMutationsOfType(io<MutType>$ mutType)
- (+)getValue(s$ key)
                                                                        g1, ... (o<GEType>$)
- (f)relatedness(o<Ind> individuals)
                                                                         i1, ... (o<IntType>$)
+ (void)setSpatialPosition(f position)
                                                                        m1, ... (o<MutType>$)
- (void)setValue(s$ key, + value)
                                                                        p1, ... (o<Subpop>$)
- (f$)sumOfMutationsOfType(io<MutType>$ mutType)
                                                                        s1, ... (o<SEBlock>$)
                                                                        self (o<SEBlock>$)
- (o<Mut>)uniqueMutationsOfType(io<MutType>$ mutType)
```

```
SLiMSim:
                                                                                SLiMgui quick help:
chromosome => (o<Chromosome>$)
                                                                                opt-click on keyword
chromosomeType => (s$)
dimensionality => (s$)
dominanceCoeffX <-> (f$)
                                                                                 Code completion:
generation <-> (i$)
                                                                                   escape (ESC) or
genomicElementTypes => (o<GEType>)
                                                                                  cmd-period (\mathbb{x}.)
inSLiMgui => (l$) (deprecated in SLiM 3.2.1)
interactionTypes => (o<IntType>)
modelType => (s$)
mutationTypes => (o<MutType>)
mutations => (o<Mut>)
nucleotideBased => (l$) (nuc)
periodicity => (string$)
scriptBlocks => (o<SEBlock>)
sexEnabled => (l$)
subpopulations => (o<Subpop>)
substitutions => (o<Substitution>)
tag <-> (i$)
- (o<Subpop>$)addSubpop(is$ subpopID, i$ size, [f$ sexRatio])
- (o<Subpop>$)addSubpopSplit(is$ subpopID, i$ size, io<Subpop>$ sourceSubpop,
      [f$ sexRatio]) (WF)
- (i$)countOfMutationsOfType(io<MutType>$ mutType)
  (void)deregisterScriptBlock(io<SEBlock> scriptBlocks)
  (+)getValue(s$ key)
  (i)mutationCounts(No<Subpop> subpops, [No<Mut> mutations])
  (f)mutationFrequencies(No<Subpop> subpops, [No<Mut> mutations])
  (o<Mut>)mutationsOfType(io<MutType>$ mutType)
  (void)outputFixedMutations([Ns$ filePath], [l$ append])
  (void)outputFull([Ns$ filePath], [l$ binary], [l$ append], [l$ spatialPositions], [l$ ages])
(void)outputMutations(o<Mut> mutations, [Ns$ filePath], [l$ append])
  (void)outputUsage(void)
- (i$)readFromPopulationFile(s$ filePath)
- (void)recalculateFitness([Ni$ generation])
 (o<SEBlock>$)register[Early/Late]Event(Nis$ id, s$ source, [Ni$ start], [Ni$ end])
  (o<SEBlock>$)registerFitnessCallback(Nis$ id, s$ source, Nio<MutType>$ mutType,
      [Nio<Subpop>$ subpop], [Ni$ start], [Ni$ end])
  (o<SEBlock>$)registerInteractionCallback(Nis$ id, s$ source, io<IntType>$ intType,
      [Nio<Subpop>$ subpop], [Ni$ start], [Ni$ end])
  (o<SEBlock>$)register[MateChoice (WF)/ModifyChild/Recombination]Callback(Nis$ id, s$ source,
      [Nio<Subpop>$ subpop], [Ni$ start], [Ni$ end])
  (o<SEBlock>$)registerMutationCallback(Nis$ id, s$ source, [Nio<MutType>$ mutType],
      [Nio<Subpop>$ subpop], [Ni$ start], [Ni$ end])
  (o<SEBlock>$)registerReproductionCallback(Nis$ id, s$ source, [Nio<Subpop>$ subpop],
      [Ns$ sex], [Ni$ start], [Ni$ end]) (nonWF)
  (o<SEBlock>$)rescheduleScriptBlock(o<SEBlock>$ block, [Ni$ start], [Ni$ end],
      [Ni generations])
- (void)setValue(s$ key, + value)
 (void)simulationFinished(void)
- (l$)treeSeqCoalesced(void)
  (void)treeSeqOutput(s$ path, [l$ simplify], [l$ includeModel])
  (void)treeSegRememberIndividuals(o<Ind> individuals)
  (void)treeSeqSimplify(void)
SLiM built-in functions:
- (s)codonsToAminoAcids(i codons, [l$ long], [l$ paste]) (nuc)
- (is)codonsToNucleotides(i codons, [s$ format]) (nuc)
- (f)mm16To256(f mutationMatrix) (nuc)
- (f)mmJukesCantor(f$ alpha) (nuc)
- (f)mmKimura(f$ alpha, f$ beta) (nuc)
- (i)nucleotideCounts(is sequence) (nuc)
- (f)nucleotideFrequencies(is sequence) (nuc)
(i)nucleotidesToCodons(is sequence) (nuc)
- (is)randomNucleotides(i$ length, [Nif basis], [s$ format]) (nuc)
```

```
Substitution:
Mutation (Mut):
                                                                id => (i\$)
id \Rightarrow (i\$)
mutationType => (o<MutType>$)
                                                                fixationTime => (i$)
nucleotide <-> (s$) (nuc)
                                                                mutationType => (o<MutType>$)
nucleotideValue <-> (i$) (nuc)
                                                                nucleotide <-> (s$) (nuc)
originGeneration => (i$)
                                                                nucleotideValue <-> (i$) (nuc)
position => (i$)
                                                                originGeneration => (i$)
selectionCoeff => (f$)
                                                                position => (i$)
subpopID <-> (i$)
                                                                selectionCoeff => (f$)
tag <-> (i$)
                                                                subpopID <-> (i$)
                                                                tag <-> (i$)
- (+)getValue(s$ key)
                                                               - (+)getValue(s$ key)
- (void)setMutationType(io<MutType>$ mutType)
- (void)setSelectionCoeff(f$ selectionCoeff)
                                                                - (void)setValue(s$ key, + value)
- (void)setValue(s$ key, + value)
Subpopulation (Subpop):
                                                                 Fitness effects of mutations:
cloningRate => (f) (WF)
firstMaleIndex => (i$)
                                                                  no mutation present
fitnessScaling <-> (f$)
                                                                                       1 + h*s
                                                                  heterozygote
genomes => (o<Genome>)
                                                                  homozygote
                                                                                         1 + s
id \Rightarrow (i\$)
immigrantSubpopFractions => (f) (WF)
immigrantSubpopIDs => (i) (WF)
                                                                 s = mut.selectionCoeff
individualCount => (i$)
                                                                 h = mutType.dominanceCoeff
individuals => (o<Ind>)
selfingRate => (f$) (WF)
sexRatio => (f$) (WF)
spatialBounds => (f)
tag <-> (i\$)
- (No<Ind>$)addCloned(o<Ind>$ parent) (nonWF)
- (No<Ind>$)addCrossed(o<Ind>$ parent1, o<Ind>$ parent2, [Nfs$ sex]) (nonWF)
- (No<Ind>$)addEmpty([Nfs$ sex]) (nonWF)
- (No<Ind>$)addRecombinant(No<Genome>$ strand1, No<Genome>$ strand2, Ni breaks1,
     No<Genome>$ strand3, No<Genome>$ strand4, Ni breaks2, [Nfs$ sex]) (nonWF)
- (No<Ind>$)addSelfed(o<Ind>$ parent) (nonWF)
- (f)cachedFitness(Ni indices)
  (void)configureDisplay([Nf center], [Nf$ scale], [Ns$ color])
  (void)defineSpatialMap(s$ name, s$ spatiality, Ni gridSize, n values, [l$ interpolate],
      [Nif valueRange], [Ns colors])
  (+)getValue(s$ key)
  (void)outputMSSample(i$ sampleSize, [l$ replace], [s$ requestedSex],
      [Ns$ filePath], [l$ append], [l$ filterMonomorphic])
  (void)outputSample(i$ sampleSize, [l$ replace], [s$ requestedSex],
      [Ns$ filePath], [l$ append])
  (void)outputVCFSample(i$ sampleSize, [l$ replace], [s$ requestedSex],
      [l$ outputMultiallelics], [Ns$ filePath], [l$ append])
- (l)pointInBounds(f point)
  (f)point[Periodic|Reflected|Stopped](f point)
- (f)pointUniform([i$ n])
- (void) removeSubpopulation(void) (nonWF)
- (void)sampleIndividuals(i$ size, [l$ replace], [No<Ind>$ exclude], [Ns$ sex], [Ni$ tag],
      [Ni$ minAge], [Ni$ maxAge], [Nl$ migrant])
- (void)setCloningRate(n rate) (WF)
- (void)setMigrationRates(io<Subpop> sourceSubpops, n rates) (WF)
- (void)setSelfingRate(n$ rate) (WF)
- (void)setSexRatio(f$ sexRatio) (WF)
- (void)setSpatialBounds(n bounds)
- (void)setSubpopulationSize(i$ size) (WF)
- (void)setValue(s$ key, + value)
- (s)spatialMapColor(s$ name, n value)
- (f)spatialMapValue(s$ name, f point)
- (void)subsetIndividuals([No<Ind>$ exclude], [Ns$ sex], [Ni$ tag], [Ni$ minAge],
      [Ni$ maxAge], [Nl$ migrant])
- (void)takeMigrants(o<Ind> migrants) (nonWF)
```

```
Genome:
genomePedigreeID => (i$)
genomeType => (s$)
individual => (o<Ind>$)
isNullGenome => (l$)
mutations => (o<Mut>)
tag <-> (i\$)
+ (void)addMutations(o<Mut> mutations)
+ (o<Mut>)addNewDrawnMutation(io<MutType> mutationType, i position, [Ni originGeneration],
      [Nio<Subpop> originSubpop], [Nis nucleotide])
+ (o<Mut>)addNewMutation(io<MutType> mutationType, n selectionCoeff, i position,
      [Ni originGeneration], [Nio<Subpop> originSubpop], [Nis nucleotide])
- (l$)containsMarkerMutation(io<MutType>$ mutType, i$ position, [l$ returnMutation])
- (l)containsMutations(o<Mut> mutations)
- (i$)countOfMutationsOfType(io<MutType>$ mutType)
- (o<Mut>)mutationsOfType(io<MutType>$ mutType)
- (is)nucleotides([Ni$ start], [Ni$ end], [s$ format])
+ (void)output([Ns$ filePath], [l$ append])
+ (void)outputMS([Ns$ filePath], [l$ append], [l$ filterMonomorphic])
+ (void)outputVCF([Ns$ filePath], [l$ outputMultiallelics], [l$ append])
- (i)positionsOfMutationsOfType(io<MutType>$ mutType)
+ (o<Mut>)readFromMS(s$ filePath, io<MutType>$ mutationType)
+ (o<Mut>)readFromVCF(s$ filePath, [Nio<MutType>$ mutationType])
+ (void)removeMutations([No<Mut> mutations], [l$ substitute])
- (f$)sumOfMutationsOfType(io<MutType>$ mutType)
Chromosome:
colorSubstitution <-> (s$)
geneConversionEnabled => (l$)
geneConversionGCBias => (f$)
geneConversionNonCrossoverFraction => (f$)
geneConversionMeanLength => (f$)
geneConversionSimpleConversionFraction => (f$)
genomicElements => (o<GElement>)
hotspotEndPositions[F[M] => (i) (nuc)
hotspotMultipliers[F|M] => (f) (nuc)
lastPosition => (i$)
mutationEndPositions[F|M] => (i)
mutationRates[F|M] => (f)
overallMutationRate[F|M] => (f$)
overallRecombinationRate[F|M] => (f$)
recombinationEndPositions[F|M] => (i)
recombinationRates[F|M] => (f)
tag <-> (i\$)
- (is)ancestralNucleotides([Ni$ start], [Ni$ end], [s$ format]) (nuc)
- (integer)drawBreakpoints([No<Ind>$ parent], [Ni$ n])
- (integer$)setAncestralNucleotides(is sequence) (nuc)
- (void)setGeneConversion(n$ nonCrossoverFraction, n$ meanLength,
      n$ simpleConversionFraction, [n$ bias])
- (void)setHotspotMap(n multipliers, [Ni ends], [s$ sex]) (nuc)
- (void)setMutationRate(n rates, [Ni ends], [s$ sex])
- (void)setRecombinationRate(n rates, [Ni ends], [s$ sex])
SLiMEidosBlock (SEBlock):
active <-> (i$)
end \Rightarrow (i$)
id \Rightarrow (i\$)
source => (s$)
start => (i$)
tag <-> (i$)
type \Rightarrow (s$)
```

```
GenomicElementType (GEType):
color <-> (s$)
id \Rightarrow (i\$)
mutationFractions => (f)
mutationMatrix => (f) (nuc)
mutationTypes => (o<MutType>)
tag <-> (i$)
- (+)getValue(s$ key)
- (void)setMutationFractions(io<MutType> mutationTypes, n proportions)
– (void)setMutationMatrix(f mutationMatrix) (nuc)
- (void)setValue(s$ key, + value)
GenomicElement (GElement):
endPosition => (i$)
genomicElementType => (o<GEType>$)
startPosition => (i$)
tag <-> (i$)
- (void)setGenomicElementType(io<GEType>$ genomicElementType)
MutationType (MutType):
color <-> (s$)
colorSubstitution <-> (s$)
convertToSubstitution <-> (l$)
distributionParams => (f)
distributionType => (s$)
dominanceCoeff <-> (f$)
id => (i\$)
mutationStackGroup <-> (i$)
mutationStackPolicy <-> (s$)
nucleotideBased => (l$) (nuc)
tag \leftarrow > (i\$)
- (float)drawSelectionCoefficient(i$ n)
- (+)getValue(s$ key)
- (void)setDistribution(s$ distType, ...)
- (void)setValue(s$ key, + value)
InteractionType (IntType):
id \Rightarrow (i\$)
maxDistance <-> (f$)
reciprocal => (l$)
sexSegregation => (s$)
spatiality => (s$)
tag <-> (i$)
- (f)distance(o<Ind> individuals1, [No<Ind> individuals2])
- (f)distanceToPoint(o<Ind> individuals1, f point)
- (o<Ind>)drawByStrength(o<Ind>$ individual, [i$ count])
- (void)evaluate([No<Subpop> subpops], [l$ immediate])
- (+)getValue(s$ key)
- (i)interactingNeighborCount(o<Ind> individuals)
- (f)interactionDistance(o<Ind> individuals1, [No<Ind> individuals2])
  (o<Ind>)nearestInteractingNeighbors(o<Ind>$ individual, [i$ count])
  (o<Ind>)nearestNeighbors(o<Ind>$ individual, [i$ count])
  (o<Ind>)nearestNeighborsOfPoint(o<Subpop>$ subpop, f point, [i$ count])
  (void)setInteractionFunction(s$ functionType, ...)
  (void)setValue(s$ key, + value)
  (f)strength(o<Ind>$ receiver, [No<Ind> exerters])
  (f)totalOfNeighborStrengths(o<Ind> individuals)
  (void)unevaluate(void)
SLiMgui: (in SLiMgui only)
pid \Rightarrow (i\$)
- (void)openDocument(s$ filePath)
(void)pauseExecution(void)
```