

# Mutations in Retroviral Genes Associated with Drug Resistance

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## INTRODUCTION

Drug resistance is the inevitable consequence of incomplete suppression of HIV replication. The rapid replication rate of HIV and its inherent genetic variation have led to the identification of many HIV variants that exhibit altered drug susceptibility. The growing number of drug resistance mutations listed in this revised table stands as a testimony to the genetic flexibility of HIV. This updated table lists 148 mutations, of which 45 occur in protease, 70 in reverse transcriptase, and 33 in envelope. Although the tables are quite comprehensive, the reader should be reminded that the mutations described are predominantly found in clade B virus and not in other HIV genotypes. The revised table also includes drug resistance mutations that have been identified for SIV and FIV.

In the table the phrase "Enzyme resist." refers to inhibition assays done just with a mutated enzyme. Instead of introducing the mutations into a virus and testing the susceptibility of the mutant virus to a drug, researchers introduce the mutation(s) into the enzyme and determine their effect by running enzyme activity assays. This type of susceptibility testing does not take into account changes in other viral proteins (like gag) that would also help confer resistance, which is the reason for distinguishing enzyme resistance from whole virus resistance.

All of the information contained in these printed tables is also available in a searchable database located at our Web site [http://204.121.6.64:581/Resist\\_DB/](http://204.121.6.64:581/Resist_DB/). There is also a facility at that site for submitting new information regarding drug resistance mutations to the database.

## ACKNOWLEDGMENTS

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**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change   | Class of Drug                  | Compound         | In vitro vivo -resistance | -Fold (-fold)   | Cross-resist  | Comments                     | Refs |
|-------------------|----------------|--------------------------------|------------------|---------------------------|---|---|------------------------------|------|
| M 41 L            | ATG to TTG/CTG | Nucleoside RTI                 | AZT              | ? Y 4                     |   | M41L/T215Y: 60-70-fold; M41L/D67N/K70R/T215Y: 180-fold.   | Larder89, Larder91, Kellam92 |      |
| A 62 V            | GCC to GTC     | Multiple Nucleoside Resistance |                  | N Y Nil                   |   | A62V alone has no effect, but in combination with mutations at 75, 77, 116, 151 causes multi NRTI resistance. | Iversen96, Shirasaka95       |      |
| K 65 R            | AAA to AGA     | Nucleoside RTI                 | 1592U89          | Y N 3                     |   | K65R/L74V: 3.6-fold; K65R/M184V: 7-fold; K65R/L74V/M184V: 10.2-fold   | Tisdale97                    |      |
| K 65 R            | AAA to AGA     | Nucleoside RTI                 | ddC              | Y Y 4-10                  | ddC; PMEA; 3TC5)  | Infrequently observed in patients receiving ddI or ddC  | Zhang94, Gu94                |      |
| K 65 R            | AAA to AGA     | Nucleoside RTI                 | ddl              | Y Y 4-10                  |   | K65R/M184V: 4.2-fold.   | Zhang94                      |      |
| K 65 R            | AAA to AGA     | Nucleoside RTI                 | dOTC (BCH-10652) | Y ?                       |   |   | Rando99                      |      |
| K 65 R            | AAA to AGA     | Nucleoside RTI                 | DXG              | Y ? 8                     | other dioxolane derivatives                                 | Reverses AZT resistance in D67N/K70R/T215Y/K219Q background   | Mellors96                    |      |
| K 65 R            | AAA to AGA     | Nucleoside RTI                 | PMEA             | Y N 10-25                 |   |   | Gu95, Fol96                  |      |
| K 65 R            | AAA to AGA     | Nucleoside RTI                 | PMPA             | Y ? 3.5                   |   | D67N/K70R/T215Y/K219Q: 120-fold; M41L/D67N/K70R/T215Y: 180-fold.  | Cherrington97                |      |
| D 67 N            | GAC to AAC     | Nucleoside RTI                 | AZT              | Y Y                       |   | Frequently associated with other multidrug resistance mutations V75I, F77L, F116Y and Q151M.                  | Larder89, Larder91, Kellam92 |      |
| S 68 G            | AGT to GGT     | Multiple Nucleoside            |                  | Y ?                       |   | Seen in one patient on 3TC + d4T combination therapy.   | Schmit98                     |      |
| T 69 A            | ACT to GCT     | Multiple Nucleoside            | 3TC + d4T        | ? Y                       | Confers >4-fold resistance to: AZT, ddI, ddC, 3TC and PMEA. | Insertion mutation. Seen in heavily treated patients.   | Lawrence99                   |      |
| T 69 D            | ACT to GAT     | Multiple Nucleoside            | AZT + 3TC        | ? Y                       |   | Seen in one patient on AZT + 3TC combination therapy.   | Winters98                    |      |
| T 69 D            | ACT to GAT     | Nucleoside RTI                 | ddC              | N Y 5                     |   |   | Fitzgibbon92                 |      |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change         | Class of Drug       | Compound          | In vitro | In vivo | -Fold -resistance   | (-fold) | Cross-resist | Comments   | Refs       |
|-------------------|----------------------|---------------------|-------------------|----------|---------|---|---------|--------------|--|------------|
| T 69 N            | ACT to AAT           | Multiple Nucleoside | 3TC + d4T         | ?        | Y       |   |         |              | Seen in two patients on 3TC + d4T combination therapy. | Lawrence99 |
| T 69 S +          | ACT to AGT + AGA GCA | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to: AZT, ddI, ddC, 3TC and PMEA. |         |              | Insertion mutation. Seen in heavily treated patients.  | Winters98  |
| T 69 S + EA       | ACT to AGT + AGA GCA | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to: AZT, ddI, ddC, 3TC and PMEA. |         |              | Insertion mutation. Seen in heavily treated patients.  | Winters98  |
| T 69 S + SA       | ACT to AGC + AGC GCT | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to: AZT, ddI, ddC, 3TC and PMEA. |         |              | Insertion mutation. Seen in heavily treated patients.  | Winters98  |
| T 69 S + SA       | ACT to TCT + AGT GCT | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to: AZT, ddI, ddC and PMEA.      |         |              | Insertion mutation. Seen in heavily treated patients.  | Winters98  |
| T 69 S + SA       | ACT to AGT + AGC GCT | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to: AZT, ddI, ddC and PMEA.      |         |              | Insertion mutation. Seen in heavily treated patients.  | Winters98  |
| T 69 S + SG       | ACT to AGT + AGT GGT | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to: AZT, ddI, ddC and PMEA.      |         |              | Insertion mutation. Seen in heavily treated patients.  | Winters98  |
| T 69 S + SG       | ACT to AGT + AGT GGT | HIV-1 Specific RTI  | ddI + hydroxyurea | ?        | Y       |   |         |              | Insertion mutation. Seen in one patient.               | DeAntoni97 |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change     | Class of Drug       | Compound          | In vitro | In vivo | -Fold resistance  | Cross-resist (-fold)                        | Comments  | Refs                            |
|-------------------|------------------|---------------------|-------------------|----------|---------|---|---|---|---------------------------------|
| T 69 S + SS       | ACT to TCT + AGC | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to:<br>AZT, ddI, ddC,<br>3TC and PMEA. |   | Insertion mutation. Seen in heavily treated patients. | Winters98                       |
| T 69 S + SS       | ACT to TCT + AGT | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to:<br>AZT, ddI, ddC,<br>3TC and PMEA. |   | Insertion mutation. Seen in heavily treated patients. | Winters98                       |
| T 69 S + SS       | ACT to AGT + AGT | HIV-1 Specific RTI  | ddI + hydroxyurea | ?        | Y       |   |   | Insertion mutation. Seen in one patient.              | DeAnton97                       |
| T 69 S + TS       | ACT to TCT + ACC | Multiple Nucleoside |                   | ?        | Y       | Confers >4-fold resistance to:<br>AZT, ddI, ddC,<br>3TC and PMEA. |   | Insertion mutation. Seen in heavily treated patients. | Winters98                       |
| K 70 E            | AAA to GAA       | Nucleoside RTI      | PMEA              | Y        | Y       | 9   | 3TC (7);<br>PFA: 2-fold hypersusceptibility |   | Cherrington96, Mulato97         |
| K 70 R            | AAA to AGA       | Nucleoside RTI      | AZT               | Y        | Y       |   |   | D67N/K70R/T215Y/K219Q: 120-fold                       | Larder89, Larder91,<br>Kellam92 |
| K 70 S            | AAA to AGA       | Multiple Nucleoside | ddI + d4T         | ?        | Y       |   |   | Seen in one patient on ddC + d4T combination therapy. |                                 |
| L 74 I            | TTA to ATA       | HIV-1 Specific RTI  | HBY 097           | Y        | ?       |   |   | K65RL74V: 3.6-fold;                                   | Klein96                         |
| L 74 V            | TTA to GTA       | Nucleoside RTI      | 1592U89           | Y        | N       | 4   |   | K65RL74V/M184V: 10.2-fold                             | Tisdale97                       |
| L 74 V            | TTA to GTA       | Nucleoside RTI      | ddI               | N        | Y       | 5-10  | ddC (4)                                     | Can reverse effect of T215Y AZT resistance mutation   | StClair91                       |
| L 74 V            | TTA to GTA       | Nucleoside RTI      | DXG               | Y        | ?       | 4   |   |   | Mellors96                       |
| L 74 V            | TTA to GTA       | HIV-1 Specific RTI  | HBY 097           | Y        | ?       |   |   |   | Klein96                         |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon      | Class of Drug                  | Compound        | In vitro | In vivo | -Fold -resistance (-fold) | Cross-resist Comments   | Refs                                     |
|-------------------|------------|--------------------------------|-----------------|----------|---------|---------------------------|---|--|
| V 75 I            | GTA to ATA | Multiple Nucleoside Resistance | Nil             | N        | Y       | Nil                       | V75I alone has no effect, but in combination with mutations at 62, 77, 116, 151 causes multi NRTI resistance. | Iversen96, Shirasaka95                   |
| V 75 I            | GTA to TTA | HIV-1 Specific RTI             | HY 097          | Y        | ?       |                           | Compensates for negative effect of G190E mutation on RT activity  | Klein96                                  |
| V 75 L            | GTA to TTA | HIV-1 Specific RTI             | HY 097          | Y        | ?       |                           |   |  |
| V 75 M            | GTA to ATG | Multiple Nucleoside            | ddC + d4T       | ?        | Y       |                           | Seen in one patient on ddC + d4T combination therapy.   | Klein96<br>Lawrence99                    |
| V 75 T            | GTA to ACA | Nucleoside RTI                 | d4T             | Y        | Y       | 7                         | Observed with d4T selection in vitro, rarely in patients receiving d4T  | Lacey94, Schinazi96                      |
| F 77 L            | TTC to CTC | Multiple Nucleoside Resistance | Nil             | N        | Y       | Nil                       | F77L alone has no effect, but in combination with mutations at 62, 75, 116, 151 causes multi NRTI resistance. | Iversen96, Shirasaka95                   |
| W 88 G            | TGG to GGG | Pyrophosphate Analogue RTI     | Foscarnet (PFA) | Y        | Y       | 5                         | Hypersusceptibility Observed after selection with AZT to AZT and PFA; suppresses effects of AZT mutations     | Mellors95, Tachedjian95,<br>Tachedjian96 |
| W 88 S            | TGG to TCG | Pyrophosphate Analogue RTI     | Foscarnet (PFA) | N        | Y       | 2-4                       | Wild-type susceptibility to AZT.  |  |
| E 89 G            | GAA to GGA | Pyrophosphate Analogue RTI     | Foscarnet (PFA) | Y        | N       | 14                        | Partially suppresses effects of AZT resistance mutations  | Mellors95, Tachedjian95,<br>Tachedjian96 |
| E 89 K            | GAA to GGA | Pyrophosphate Analogue RTI     | Foscarnet (PFA) | Y        | N       | > 16                      | Isolated by screening RT clones for ddGTP resistance  | Prasad91                                 |
| L 92 I            | TTA to ATA | Pyrophosphate Analogue RTI     | Foscarnet (PFA) | Y        | N       | 8                         | Suppresses effects of AZT resistance mutations  | Tachedjian95, Tachedjian96               |
| A 98 G            | GCA to GGA | HIV-1 Specific RTI             | L-697,661       | N        | Y       | 8                         | Partially suppresses effects of AZT resistance mutations  | Byrnes93                                 |
| A 98 G            | GCA to GGA | HIV-1 Specific RTI             | Nevirapine      | N        | Y       |                           |   | Richman94                                |
| L 100 I           | TTA to ATA | HIV-1 Specific RTI             | BHAP U-88204E   | Y        | ?       |                           |   | Bazarini93d, Vasudevachari92             |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid | Codon Change | Class of Drug      | Compound             | In vitro | In vivo | -Fold resistance | Cross-resist (-fold) | Comments  | Refs   |
|------------|--------------|--------------------|----------------------|----------|---------|------------------|----------------------|---|--|
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | DMP 266 (L-743,726)  | Y        | ?       | 8-11             |                      | Combinations of mutations needed for high-level resistance; L100I/V108I: 1,000-fold; L100I/V179D/Y181C: 1,000-fold                                  | Young95, Winslow96   |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | L-697,661 Nevirapine | Y        | N       | 2                |                      |   | Byrnes93   |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | TIBO R82150          | Y        | ?       | > 100            |                      |   | Richman93  |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | TIBO R82913          | Y        | ?       |                  |                      | Suppresses effects of AZT resistance mutations  | Mellors93, Balzarini93c, Byrnes93a   |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | UC-68 (638532)       | Y        | ?       | 70               |                      |   | Landry92   |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | UC-70 (638534)       | Y        | ?       | 758              |                      |   | Balzarini95  |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | UC-781               | Y        | ?       | 20               |                      |   | Buckheit95a  |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | UC-781               | Y        | ?       |                  |                      | Activity of UC-781 versus L100I, K103N, V106A, E138K, Y181C and Y188L reduced by 2-, 7-, 1.5-, 5- and 150-fold, respectively, compared to wild type | Balzarini96a, Balzarini96b   |
| L 100 I    | TTA to ATA   | HIV-1 Specific RTI | UC-84 (615985)       | Y        | ?       | > 40, > 33       |                      |   | Buckheit95a, Buckheit95b   |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | 8-Chloro-TBO R091767 | ?        | Y       |                  |                      |   | Moeremans95  |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | ADAMII               | Y        | ?       | 30               |                      |   | Cushman98  |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | Ateviridine + AZT    | ?        | Y       |                  |                      |   | Not selected for in vitro, resistance determined against a panel of mutants. Viruses with the L100I mutation show an enhanced sensitivity to ADAMII. |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | Ateviridine + AZT    | ?        | Y       |                  |                      |   | Seen in one patient on atevirdine + AZT combination therapy. Found in association with K103N.  |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | DMP 266 (L-743,726)  | Y        | ?       | 1,000            |                      |   | Demeter98  |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | L-697,661            | N        | Y       | 8                |                      |   | Young95  |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | UC-10 (645129)       | Y        | ?       | 12               |                      |   | Byrnes93   |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | UC-38 (629243)       | Y        | N       |                  |                      |   | Buckheit95a, Buckheit97  |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI | UC-57 (647014)       | Y        | ?       |                  |                      |   | Balzarini95a, Balzarini95  |
| K 101 E    | AAA to GAA   | HIV-1 Specific RTI |                      |          |         |                  |                      |   | Buckheit95a  |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound                     | In vitro | In vivo | -Fold resistance | Cross-resist (-fold)         | Comments   | Refs                 |
|-------------------|--------------|--------------------|------------------------------|----------|---------|------------------|------------------------------|--|----------------------|
| K 101 E           | AAA to GAA   | HIV-1 Specific RTI | UC-781                       | Y        | ?       | 7                | UC040 (18); Nevirapine (15)  | V108I/Y181C; 55-fold; K101E/V108I/Y181C; 500-fold.   | Buckheit97           |
| K 101 I           | AAA to ATA   | HIV-1 Specific RTI | UC-16                        | Y        | N       | 10               | K101/G141E; 10-fold          |  | Balzarini95          |
| K 101 Q           | AAA to CAA   | HIV-1 Specific RTI | Trovirdine                   | Y        | ?       |                  |                              | Found in combination with V108I  | Zhang95, Vrang93     |
| K 103 E           | AAA to GAA   | Nucleoside RTI     | BHAP U-87201E (Ateviridine)  | ?        | Y       |                  |                              | Found in association with Y181C in one patient on monotherapy. K103E, K103N and Y181C observed with monotherapy.                                     | Demeter98            |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | 8-Chloro-TIBO R091767        | ?        | Y       |                  |                              |  | Moremans95           |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | ADAMII                       | Y        | ?       | >28              |                              | Not selected for in vitro, resistance determined against a panel of mutants. Viruses with the L100I mutation show an enhanced sensitivity to ADAMII. | Cushman98            |
| K 103 N           | AAA to AAC   | Nucleoside RTI     | BHAP U-87201E (Ateviridine)  | ?        | Y       |                  |                              | Found in association with Y181C in several patients on monotherapy. Also seen in patients on ATV + AZT combination therapy.                          | Demeter98            |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | BHAP U-90152 (delavirdine)   | ?        | Y       |                  |                              | K103N/Y181C seen separately and in combination in patients   | Klein99., Klein99(2) |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | DMP 266 (L-743,726)          | Y        | Y       | 67               | Predominant mutation in vivo | Winslow96  |                      |
| K 103 N           | AAA to AAC   | Nucleoside RTI     | GW420867X                    | Y        | ?       |                  |                              | K103N/Y181C; > 1,000-fold  |                      |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | L-697,593                    | Y        | ?       | 20               |                              | K103N and Y181C most common with monotherapy   | Nurnberg91           |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | L-697,661                    | Y        | Y       | 8                |                              |  | Byrnies93, Saag93    |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | Loviride (R89439, alpha-APA) | Y        | Y       |                  |                              |  | Staszewski96a        |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | MKC442 (I-EBU)               | Y        | ?       |                  |                              | Predominant mutation in vivo   | Seki95               |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | Nevirapine                   | N        | Y       |                  |                              |  | Richman93            |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | TIBO R82913                  | Y        | ?       | > 100            |                              |  | Balzarini93d         |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | UC-10 (645129)               | Y        | N       | 5                |                              |  | Balzarini95          |
| K 103 N           | AAA to AAC   | HIV-1 Specific RTI | UC-81 (615727)               | Y        | ?       |                  |                              |  | Balzarini95, Yang97  |
| K 103 Q           | AAA to CAA   | HIV-1 Specific RTI | L-697,661                    | N        | Y       | 8                |                              |  | Saag93               |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug                    | Compound       | In vitro | In vivo | -Fold -resistance | Cross-resist (-fold)      | Comments  | Refs   |
|-------------------|--------------|----------------------------------|----------------|----------|---------|-------------------|---------------------------|---|--|
| K 103 R           | AAA to AGA   | HIV-1 Specific RTI               | MKC442 (1-EBU) | Y        | Y       | ?                 | Nevirapine; 9-chloro-TIBO | K103R/V179D; 500-fold; Found in combination with V179D or Y181C | BorrottoEsoda97<br>Zhang95, Vrang93          |
| K 103 R           | AAA to AGA   | HIV-1 Specific RTI               | Trovirdine     | Y        | ?       | ?                 |                           |   | Demeter95                                    |
| K 103 T           | AAA to ACA   | HIV-1 Specific RTI (delavirdine) | BHAP U-90152   | ?        | Y       |                   |                           |   |  |
| K 103 T           | AAA to ACA   | Nucleoside RTI                   | S-1153         | Y        | ?       |                   |                           |   | Fujiiwara98                                  |
| K 103 T           | AAA to ACA   | HIV-1 Specific RTI               | UC-42          | Y        | N       | 100               |                           |   | Balzarini95                                  |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI               | ADAMII         | Y        | ?       | 7.13              |                           |   | Cushman98                                    |
|                   |              |                                  |                |          |         |                   |                           |   |  |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI               | BHAP U-88204E  | Y        | ?       |                   |                           |   | Vasudevachari92                              |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI               | E-EBU-dM       | Y        | ?       |                   |                           |   | Balzarini93                                  |
| V 106 A           | GTA to GCA   | Nucleoside RTI                   | GW420867X      | Y        | ?       |                   |                           |   | Klein99, Kleim99(2)                          |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI               | Nevirapine     | Y        | Y       | 100               |                           |   | Richman94, Larder92, Richman93, Balzarini93d |
| V 106 A           | GTA to GCA   | Nucleoside RTI                   | S-1153         | Y        | ?       | 4.5               |                           |   | Fujiiwara98                                  |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI (Quinoxaline) | S-2720         | Y        | ?       |                   |                           |   | Pelmans97                                    |
|                   |              |                                  |                |          |         |                   |                           |   |  |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI               | TIBO R82913    | Y        | ?       | 100               |                           |   | Larder92                                     |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI               | UC-69 (646989) | Y        | ?       | ?                 |                           |   | V106AV181C; 166-fold                         |
| V 106 A           | GTA to GCA   | HIV-1 Specific RTI               | UC-82          | Y        | ?       | 13                |                           |   | Buckheit95a                                  |
|                   |              |                                  |                |          |         |                   |                           |   | Balzarini96b, Balzarini96a                   |
| V 106 I           | GTA to ATA   | HIV-1 Specific RTI               | HBY 097        |          |         |                   |                           |   | Kleim97                                      |
|                   |              |                                  |                |          |         |                   |                           |   |  |
|                   |              |                                  |                |          |         |                   |                           |   |  |

## Mutations in HIV RT that confer drug resistance, ordered by position.

| Amino Acid                      | Codon Change | Class of Drug                  | Compound                             | In vitro | In vivo | -Fold -resistance (-fold)   | Cross-resist Comments  | Refs                                   |  |  |
|---------------------------------|--------------|--------------------------------|--------------------------------------|----------|---------|---|--|--|--|--|
| V 108 I                         | GTA to ATA   | HIV-1 Specific RTI             | ADAMII                               | Y        | ?       | 6.74  | Not selected for in vitro, resistance determined against a panel of mutants. Viruses with the L100I mutation show an enhanced sensitivity to ADAMII. |  |  |  |
| V 108 I                         | GTA to ATA   | HIV-1 Specific RTI             | DMP 266 (L-743,726)                  | Y        | ?       | L100I/V108I: 1,000-fold   |  | Winslow96                              |  |  |
| V 108 I                         | GTA to GCA   | HIV-1 Specific RTI             | L-697,661                            | Y        | Y       | 4   | Byrnes93   |  |  |  |
| V 108 I                         | GTA to ATA   | HIV-1 Specific RTI             | Loviride (R89439, <i>alpha</i> -APA) | Y        | ?       | Staszewski96a   |  |  |  |  |
| V 108 I                         | GTA to GCA   | HIV-1 Specific RTI             | MKC442 (I-EBU)                       | Y        | ?       | Seki95  |  |  |  |  |
| V 108 I                         | GTA to ATA   | HIV-1 Specific RTI             | Nevirapine                           | N        | Y       | Richman93   |  |  |  |  |
| V 108 I                         | GTT to GAT   | HIV-1 Specific RTI             | TIBO R 82913                         | N        | Y       | > 100   | R82150 (> 100)   |  |  |  |
| V 108 I                         | GTA to ATA   | HIV-1 Specific RTI             | Trovirdine                           | Y        | ?       | Vandamme94a   |  |  |  |  |
| V 108 I                         | GTA to ATA   | HIV-1 Specific RTI             | UC-781                               | Y        | ?       | Zhang95   |  |  |  |  |
| Found in combination with K101Q |              |                                |                                      |          |         |   |  |  |  |  |
| Buckheit97                      |              |                                |                                      |          |         |   |  |  |  |  |
| V108I/Y181C:<br>55 fold.        |              |                                |                                      |          |         |   |  |  |  |  |
| K101E/V108I/Y181C:<br>500 fold. |              |                                |                                      |          |         |   |  |  |  |  |
| Y 115 F                         | TAT to TTT   | Nucleoside RTI                 | 1592U89                              | Y        | N       | 2   | Tisdale97<br>K65R/I/74V and/or Y115F with M184V: 10 fold; L74V/Y115F/M184V: 11-fold  |  |  |  |
| F 116 Y                         | TTT to TAT   | Multiple Nucleoside Resistance |                                      | N        | Y       | Nil   | Iversen96, Shirasaka95<br>F116Y alone has no effect, but in combination with mutations at 62, 75, 77, 151 causes multi NRTI resistance.              |  |  |  |
| P 119 S                         | CCC to TCC   | Nucleoside RTI                 | F-ddA                                | Y        | ?       | 4.6   | Found with V179D and/or L214F, which are possibly compensatory   |  |  |  |
| E 138 K                         | GAG to AAG   | HIV-1 Specific RTI             | MKC442 (I-EBU)                       | Y        | N       | Obtained in the concomitant presence of low 3TC concentrations          |  | Balzarini96c                           |  |  |
| E 138 K                         | GAG to AAG   | HIV-1 Specific RTI             | TIBO R 82913                         | Y        | ?       | Balzarini93c  |  |  |  |  |
| E 138 K                         | GAG to AAG   | HIV-1 Specific RTI             | TSAO                                 | Y        | ?       | E138A (GAG to GCG) in TSAO-naive patients confers TSAO viral resistance |  | Balzarini93a, Balzarini93b, Vandamme96 |  |  |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug                  | Compound            | In vitro       | In vivo | -Fold -resistance (-fold) | Cross-resist Comments   | Refs   |           |
|-------------------|--------------|--------------------------------|---------------------|----------------|---------|---------------------------|---|--|-----------|
| E 138 K           | GAG to AAG   | HIV-1 Specific RTI             | UC-82               | Y              | ?       | 5                         |   | Balzarini96b, Balzarini96a   |           |
| T 139 I           | ACA to ATA   | HIV-1 Specific RTI             | ADAMII              | UC-84 (615985) | Y       | ? > 100                   | TSAOs   | Balzarini95, Balzarini95b Cushman98  |           |
|                   |              |                                |                     |                | Y       | ?                         | 38  |  |           |
| T 139 I           | ACA to ATA   | HIV-1 Specific RTI             | Calanolide A        | Y              | ?       | > 70                      | Not other NNRTIs  | Buckheit95c  |           |
| G 141 E           | GGG to GAG   | HIV-1 Specific RTI             | UC-16               | Y              | N       |                           | K101I/G141E: 10-fold  | Balzarini95  |           |
| Q 151 M           | CAG to ATG   | Multiple Nucleoside Resistance |                     | N              | Y       | AZT; 10; ddI/ ddC; 5      | Pivotal multi nucleoside RTI resistance mutation (first to occur), found in association with combinations of four other mutations: A62V/ V75I/ F77L/ F116Y/Q151M; AZT 190-fold; ddI 50-fold; ddC 20-fold; d4T > 10-fold | Iversen96, Shirasaka95, Schmit96   |           |
| S 156 A           | TCA to GCA   | Pyrophosphate Analogue RTI     | Foscarnet (PFA)     | Y              | N       | 4.5                       |   | Tachdjian95  |           |
| Q 161 L           | CAA to CTA   | Pyrophosphate Analogue RTI     | Foscarnet (PFA)     | Y              | Y       | 5                         | Q161L/H208Y: 9-fold; Q161L/H208Y suppresses effects of AZT mutations  | Mellors95  |           |
| V 179 D           | GTT to GAT   | HIV-1 Specific RTI             | ADAMII              | Y              | ?       | 28                        |   | Cushman98  |           |
| V 179 D           | GTT to GAT   | HIV-1 Specific RTI             | DMP 266 (L-743,726) | Y              | ?       |                           |   | Not selected for in vitro, resistance determined against a panel of mutants. Viruses with the L100I mutation show an enhanced sensitivity to ADAMII. L100I/V179D/Y181C: 1,000-fold | Winslow96 |
| V 179 D           | GTT to GAT   | HIV-1 Specific RTI             | L-697,661           | N              | Y       | 4                         |   | Byrnes93   |           |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid | Codon Change | Class of Drug      | Compound                              | In vitro | In vivo | -Fold -resistance | Cross-resist (-fold)  | Comments   | Refs                      |
|------------|--------------|--------------------|---------------------------------------|----------|---------|-------------------|---|--|---------------------------|
| V 179 D    | GTT to GAT   | HIV-1 Specific RTI | QM96521                               | Y        | ?       | 10                | Other TDD derivative: 15-, 140-fold; 8-chloro-TIBO: 10-fold |  | Witvrouw98                |
| V 179 D    | GTT to GAT   | HIV-1 Specific RTI | TIBO R82913                           | N        | Y       | 20                | R82150 (20)   | Found in combination with K103R or Y181C; V179D/Y181C: > 1,000-fold  | Vandamme94<br>Zhang95     |
| V 179 D    | GTT to GAT   | HIV-1 Specific RTI | Trovirdine                            | Y        | ?       | 16                |   |  | Balzarini95, Balzarini96a |
| V 179 E    | GTT to GAG   | HIV-1 Specific RTI | UC-10 (645129)                        | Y        | ?       | 16                |   |  | Byrnes93                  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | L-697,661                             | N        | Y       | 8                 |   |  | Harari97                  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | 1737 (Tetrahydrophthalene derivative) | Y        | ?       | 20                |   | Y181C also confers resistance to numerous other tetrahydronaphthalene derivatives.   |                           |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | ADAMII                                | Y        | ?       | >28               |   | Not selected for in vitro, resistance determined against a panel of mutants. Viruses with the L100I mutation show an enhanced sensitivity to ADAMII. | Cushman98<br>deBethune93  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | <i>alpha</i> -APA                     | Y        | ?       |                   |   |  |                           |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | R1893 (loviride analogue)             |          |         |                   |   |  |                           |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | BHAP U-87201E (ateviridine)           | N        | Y       |                   |   | K103E, K103N and Y181C observed with monotherapy   | Demeter95, Demeter98      |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | BHAP U-88204E                         | Y        | ?       |                   |   |  | Vasudevachari92           |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | BHAP U-90152 (delavirdine)            | ?        | Y       |                   |   | K103N/Y181C seen separately and in combination in vivo   | Demeter95                 |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | BM+51.0836                            | Y        | ?       |                   |   |  | Mass93                    |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | DMP 266 (L-743,726)                   | Y        | ?       | 4                 |   | L100I/V179D/Y181C: 1,000-fold; uncommon in vivo  | Winslow96, Young95        |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | E-EBU                                 | Y        | ?       |                   |   |  | Balzarini93               |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | E-EPSeU                               | Y        | ?       | >50               |   | Y188C confers greater resistance than Y181C  | Nguyen94                  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | E-EPU                                 | Y        | ?       | >95               |   | Y188C confers greater resistance than Y181C  | Nguyen94                  |
| Y 181 C    | TAT to TGT   | Nucleoside RTI     | GW420867X                             | Y        | ?       |                   |   |  | Klein99, Kleinm99(2)      |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid | Codon Change | Class of Drug      | Compound                             | In vitro | In vivo | -Fold -resistance | (fold) | Cross-resist   | Comments | Refs   |
|------------|--------------|--------------------|--------------------------------------|----------|---------|-------------------|--------|--|----------|--|
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | L-697,593                            | Y        | ?       | > 100             |        | K103NY181C: > 1,000-fold   |          | Nurnberg91   |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | L-697,661                            | Y        | Y       | > 30              |        | K103N and Y181C most common with monotherapy                       |          | Byrnes93, Saag93   |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | Lovinide (R89439, <i>alpha</i> -APA) | ?        | Y       |                   |        |  |          | Staszewski96   |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | MKC442 (1-EBU)                       | ?        | Y       |                   |        |  |          | BorrottoEsoda97  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | Nevirapine                           | Y        | Y       | > 100             |        |  |          | Richman94, Richman91, Mellors92                                  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | NSC 648400 (E-BPTU)                  | Y        | ?       | 160               |        |  |          | Buckheit95c  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | TIBO R82913                          | Y        | ?       | > 100             |        |  |          |  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | Trovirdine                           | Y        | ?       |                   |        |  |          |  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-10 (645129)                       | Y        | ?       | 6                 |        | K103NY181C: > 1,000-fold   |          | Lander92   |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-32 (645542)                       | Y        | ?       | 38                |        | V179DY181C: > 1,000-fold; Found in combination with K103R or V179D |          | Zhang95, Vrang93   |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-38 (629243)                       | Y        | ?       | 8-149             |        | K101EY181C: 200-fold   |          | Buckheit95a  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-57 (647014)                       | Y        | ?       |                   |        |  |          | Buckheit95a  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-68 (638532)                       | Y        | ?       | 5                 |        |  |          | Buckheit95a  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-69 (646989)                       | Y        | ?       |                   |        |  |          | Buckheit95a  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-781                               | Y        | ?       | 13                |        | V106AV181C: 166-fold   |          |  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-80 (639475)                       | Y        | ?       | 18                |        | V108/Y181C: 55 fold; K101E/ V108/Y181C: 500 fold; 42               |          | Buckheit95a  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-81 (615727)                       | Y        | ?       | 53                |        |  |          | Balzarini95, Yang97  |
| Y 181 C    | TAT to TGT   | HIV-1 Specific RTI | UC-84 (615985)                       | Y        | ?       | > 118             |        |  |          | Buckheit95a  |
| Y 181 I    | TGT to ATT   | HIV-1 Specific RTI | BHAP U-88204E                        | Y        | Y       |                   |        |  |          | Balzarini94  |
| Y 181 I    | TAT to ATT   | HIV-1 Specific RTI | MKC442 (1-EBU)                       | Y        | N       | 1,000             |        |  |          | Balzarini96c   |
| Y 181 I    | TGT to ATT   | HIV-1 Specific RTI | Nevirapine                           | N        | Y       | High-level        |        |  |          | Shaw94   |
| M 184 I    | ATG to ATA   | Nucleoside RTI     | 3TC (lamivudine)                     | Y        | Y       |                   |        |  |          | Schinazi93, Tisdale93, Gao93                                     |
|            |              |                    |                                      |          |         |                   |        |  |          | Observed in one patient  |
|            |              |                    |                                      |          |         |                   |        |  |          | MI84V and MI84I can suppress effects of AZT resistance mutations |

Mutations in HIV RT that confer drug resistance, ordered by position.

| Amino Acid | Codon Change | Class of Drug      | Compound         | In vitro | In vivo | -Fold resistance | Cross-resist (-fold)   | Comments   | Refs              |
|------------|--------------|--------------------|------------------|----------|---------|------------------|--|--|-------------------|
| M 184 T    | ATG to ACG   | Nucleoside RTI     | 3TC (lamivudine) | Y        | ?       |                  |  | Reduced replication capacity and RT activity   | Larde95, Keulen96 |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | 1592U89          | Y        | N       | 3                |  | K65R/L74V and/or Y115F with M184V: 10-fold; K65R/M184V: 8-fold; L74V/M184V: 9-fold resistance; L74V/Y115F/M184V: 11-fold | Tisdale97         |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | 3TC (lamivudine) | Y        | Y       | >100             | ddI; ddC; (-)-FTC M184V and M184I can suppress effects of AZT resistance mutations; GTA seen in cell culture   | Schinaz93, Tisdale93, Gao93  |                   |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | ddC              | Y        | Y       | 2-5              |  |  | Gu92              |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | ddI              | Y        | Y       | 2-5              |  |  | Gu92              |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | (-)dOTC          | Y        | ?       | nil              |  | Rando99  |                   |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | (+)-dOTC         | Y        | ?       |                  |  | Rando99  |                   |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | dOTC (BCH-10652) | Y        | ?       |                  | K65R/M184V: 4.2-fold.  | Rando99  |                   |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | (-) -FTC         | Y        | ?       | > 100            | MI84V can suppress effects of AZT mutations  | Schinaz93, Tisdale93   |                   |
| M 184 V    | ATG to GTG   | Nucleoside RTI     | L-FddC           | Y        | ?       | > 100            |  | Schinaz95  |                   |
| Y 188 C    | TAT to TGT   | HIV-1 Specific RTI | ADAMII           | Y        | ?       | 6.07             | Not selected for in vitro, resistance determined against a panel of mutants. Viruses with the L100I mutation show an enhanced sensitivity to ADAMII. | Cushman98  |                   |
| Y 188 C    | TAT to TGT   | HIV-1 Specific RTI | E-EPSeU          | Y        | ?       | > 250            | Y188C is the predominant mutation for E-EPSeU; Y188C confers greater resistance than Y181C   | Nguyen94   |                   |
| Y 188 C    | TAT to TGT   | HIV-1 Specific RTI | E-EPU            | Y        | ?       | > 250            | Y188C confers greater resistance than Y181C  | Nguyen94   |                   |
| Y 188 C    | TAT to TGT   | HIV-1 Specific RTI | HEPT             | Y        | ?       |                  |  | Balzarini93  |                   |
| Y 188 C    | TAT to TGT   | HIV-1 Specific RTI | Nevirapine       | N        | Y       |                  |  | Richman93  |                   |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change   | Class of Drug                  | Compound                             | In vitro | In vivo | -Fold resistance | Cross-resist (-fold) | Comments   | Refs         |
|-------------------|----------------|--------------------------------|--------------------------------------|----------|---------|------------------|----------------------|--|--------------|
| Y 188 H           | TAT to CAT     | HIV-1 Specific RTI             | ADAMII                               | Y        | ?       | >128             |                      | Not selected for in vitro, resistance determined against a panel of mutants. Viruses with the L100I mutation show an enhanced sensitivity to ADAMII. | Cushman98    |
| Y 188 H           | TAT to CAT     | HIV-1 Specific RTI             | Ateviridine + AZT                    | ?        | Y       |                  |                      | Found in two patients on atevirdine + AZT combination therapy.   | Demeter98    |
| Y 188 H           | TAT to CAT     | HIV-1 Specific RTI             | TIBO R82913                          | Y        | ?       |                  |                      |  | Balzarini93c |
| Y 188 H/L         | TAT to CAT/CCT | HIV-1 Specific RTI             | Loviride (R89439, <i>alpha</i> -APA) | ?        | Y       |                  |                      |  | Staszewski96 |
| Y 188 L           | TAT to TTA     | HIV-1 Specific RTI             | DMP 266 (L-743,726)                  | Y        | ?       | 1,000            |                      |  | Winslow96    |
| Y 188 L           | TAT to TTA     | HIV-1 Specific RTI             | TIBO R82913                          | N        | Y       |                  |                      |  | Vandamme94   |
| V 189 I           | GTA to ATA     | HIV-1 Specific RTI             | HBY 097                              | Y        | ?       | 2                |                      |  | Klein96      |
| G 190 A           | GGA to GCA     | HIV-1 Specific RTI             | Loviride (R89439, <i>alpha</i> -APA) | ?        | Y       |                  |                      |  | Moeremans95  |
| G 190 A           | GGA to GCA     | HIV-1 Specific RTI             | Nevirapine                           | N        | Y       |                  |                      |  | Richman94    |
| G 190 E           | GGA to GAA     | HIV-1 Specific RTI             | AAP-BHAP (U-104489)                  | Y        | ?       | >100             |                      |  | Ohmsted96    |
| G 190 E           | GGA to GAA     | HIV-1 Specific RTI             | GW420867X                            | Y        | ?       |                  |                      |  |              |
| G 190 E           | GGA to GAA     | HIV-1 Specific RTI             | HBY 097                              | Y        | ?       |                  |                      |  |              |
| G 190 E           | GGA to GAA     | HIV-1 Specific RTI             | S-2720                               | Y        | ?       |                  |                      |  |              |
| G 190 E           | GGA to GAA     | HIV-1 Specific RTI             | UC-38 (629243)                       | Y        | N       |                  |                      |  |              |
| G 190 Q           | GGA to CAA     | HIV-1 Specific RTI             | HBY 097                              | Y        | ?       |                  |                      |  |              |
| G 190 T           | GGA to ACA     | HIV-1 Specific RTI             | HBY 097                              | Y        | ?       |                  |                      |  |              |
| H 208 Y           | CAT to TAT     | Multiple Nucleoside Resistance | AZT + 3TC                            | ?        | Y       |                  |                      |  |              |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug                  | Compound        | In vitro | In vivo | -Fold resistance | Cross-resist (-fold)   | Comments   | Refs              |
|-------------------|--------------|--------------------------------|-----------------|----------|---------|------------------|--|--|-------------------|
| H 208 Y           | CAT to TAT   | Pyrophosphate Analogue RTI     | Foscarnet (PFA) | Y        | Y       | 2                |  | Q161L/H208Y: 9-fold; increased susceptibility to AZT 100-fold, nevirapine (20-fold) and TIBO R82150 (30-fold); Q161L/H208Y suppresses effects of AZT mutations | Mellors95         |
| L 210 W           | TTG to TGG   | Nucleoside RTI                 | AZT             | Y        | Y       |                  | 210W/215Y: 42-fold 41L/210W/215Y; 49-fold 41L/67N/70R/210W/215Y: 366-fold Mutation arises after prolonged AZT therapy.                                 | Gurusinghe95, Harrigan96, Hooker96   |                   |
| R 211 K           | AGG to AAG   | Multiple Nucleoside Resistance | AZT + 3TC       | ?        | Y       |                  |  | Polymorphism facilitating AZT+3TC dual resistance in association with M184V and other AZT resistance mutations.  | Kemp98            |
| L 214 F           | CTT to TTT   | Multiple Nucleoside Resistance | AZT + 3TC       | ?        | Y       |                  |  | Polymorphism facilitating AZT+3TC dual resistance in association with M184V and other AZT resistance mutations.  | Kemp98, Stuyver97 |
| T 215 F           | ACC to TTC   | Nucleoside RTI                 | AZT             | ?        | Y       |                  | K67N/K70R/T215Y/K219Q: 120-fold M41L/T215Y: 60-70-fold;  | Larde91, Larde91, Kellam92   |                   |
| T 215 Y           | ACC to TAC   | Nucleoside RTI                 | AZT             | Y        | Y       |                  | K67N/K70R/T215Y/K219Q: 120-fold. Effect of T215Y is reversed by a ddI mutation (L74V), NNRTI mutations (L100I;Y181C) or (-)FTC/3TC mutations (M184I/V) | Larde91, Larde91, Kellam92   |                   |
| Y 215 C           | TTC to TGC   | Nucleoside RTI                 | ddC             | N        | Y       | 4                | Arises on background of T215Y AZT resistance   | Slade93  |                   |
| K 219 E           | AAA to GAA   | Nucleoside RTI                 | AZT             | Y        | N       |                  |  | Larde91, Kellam92  |                   |
| K 219 Q           | AAA to CAA   | Nucleoside RTI                 | AZT             | ?        | Y       |                  | K67N/K70R/T215Y/K219Q: 120-fold  | Larde91, Kellam92  |                   |
| K 219 R           | AAA to AGA   | Multiple Nucleoside            | 3TC + d4T       | ?        | Y       |                  | Seen in two patient on 3TC + d4T combination therapy.  | Lawrence99   |                   |
| K 219 R           | AAA to AGA   | Multiple Nucleoside            | AZT + 3TC       | ?        | Y       |                  | Seen in two patient on AZT + 3TC combination therapy.  | Lawrence99   |                   |

**Mutations in HIV RT that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug                    | Compound          | In vitro | In vivo | -Fold -resistance (-fold) | Cross-resist Comments  | Refs                 |
|-------------------|--------------|----------------------------------|-------------------|----------|---------|---------------------------|--|----------------------|
| K 219 W           | AAA to TGG   | Multiple Nucleoside              | ddC + d4T         | ?        | Y       |                           | Seen in one patient on ddC + d4T combination therapy.  | Lawrence99           |
| P 225 H           | CCT to CAT   | HIV-1 Specific RTI (Quinoxaline) | S-2720            | Y        | ?       | 4.0                       | MKC-442 (5.7); P225H follows V106A. Also seen with L101I and Y181C. Double and triple mutants highly resistant to other NNRTI's, including MKC442. The presence of P225H in a V106A background restores sensitivity to BHAP U-90152. | Pelmans97, Pelmans98 |
| F 227 L           | TTA to CTC   | Nucleoside RTI                   | S-1153            | Y        | ?       | nil                       | V106A + F227L: 387-fold. This mutation confers hypersensitivity to delavirdine.  | Fujiwara98           |
| F 227 L           | TTA to CTC   | HIV-1 Specific RTI               | UC-781            | Y        | ?       |                           | V106AF227L: 10-fold. Found with V106A, K101I, Y181C and L101I. Appears in a V106A background following dose-escalating UC-781 treatment.   | Balzarini98          |
| V 233 E           | GAA to GTA   | HIV-1 Specific RTI               | Ateviridine + AZT | N        | Y       |                           | Seen in 1 patient. K101E, Y188H and K238T also seen in patients on ATV/AZT combination therapy.  | Demeter98            |
| L 234 I           | CTC to ATC   | Nucleoside RTI                   | S-1153            | Y        | ?       | 22-fold                   | This mutation confers hypersensitivity to Loviride.  | Fujiwara98           |
| P 236 L           | CCT to CTT   | HIV-1 Specific RTI (ateviridine) | BHAP U-87201E     | Y        | N       |                           |  | Dueweke93            |
| P 236 L           | CCT to CTT   | HIV-1 Specific RTI (delavirdine) | BHAP U-90152      | Y        | Y       |                           | Sensitizes RT 10-fold to nevirapine, TIBO R82913 and L-697,661   | Dueweke93            |
| K 238 T           | AAA to ACA   |                                  | HEPT              | Y        | ?       |                           |  | Buckheit95c          |
|                   |              |                                  | Ateviridine + AZT | N        | Y       |                           | Seen in 1 patient. K101E, K103N, Y188H, and V233E also observed with ATV/AZT combination therapy.  | Demeter98            |
| G 333 D           | GGC to GAC   | Multiple Nucleosides             | AZT+3TC           | Y        | Y       |                           | Facilitates dual resistance to AZT+3TC in association with M184V and standard AZT resistance mutations.  | Kemp98               |
| G 333 E           | GGC to GAG   | Multiple Nucleosides             | AZT + 3TC         | Y        | Y       |                           | Facilitates dual resistance to AZT+3TC in association with M184V and standard AZT resistance mutations.  | Kemp98               |

## Mutations in HIV Protease that confer drug resistance, ordered by position.

| Amino Acid Change | Codon Change | Class of Drug      | Compound        | In vitro | In vivo | -Fold -resistance | Cross-resist (-fold) | Comments  | Refs                       |
|-------------------|--------------|--------------------|-----------------|----------|---------|-------------------|----------------------|---|----------------------------|
| R 8 K             | CGA to AAA   | Protease Inhibitor | A-77003         | Y        | ?       | 10                |                      | R8K/ M46I/ G48V: 20-fold  | Ho94, Tisdale94            |
| R 8 Q             | CGA to CAA   | Protease Inhibitor | A-77003         | Y        | ?       | 10                |                      | M46I improves replication competency of R8Q mutant  | Ho94, Kaplan94             |
| L 10 F            | CTC to TTC   | Protease Inhibitor | ABT-378         | Y        | ?       |                   |                      | Passage 17 virus: I84V/L10F/M46I: 4 fold, I84V/ L10F/ ritonavir: 21-fold; M46I/T91S: 12 fold, I84V// L10F/ saquinavir, 4-fold   | Carillo98                  |
|                   |              |                    |                 |          |         |                   |                      | M46I/ T91S/ V32I/ V32L/ I47V: 46 fold, Passage 17 virus: I84V/ L10F/ M46I/ T91S/ V32L/ I47V/ V47A/ G16E/ H69Y: 338 fold (in presence of p7/p1 (AN/F to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation). |                            |
| L 10 F            | CTC to TTC   | Protease Inhibitor | BILA 2185 BS    | Y        | ?       |                   |                      | BILA 1906 BS (360)  | Croteau97                  |
|                   |              |                    |                 |          |         |                   |                      | L10F/ L23I/ V32I/ M46I/ I47V/ I54M/ A71V/ I84V: 1,500-fold. Associated Gag mutations: p1'/p6 cleavage site (L to F (CTT to TTT at P1'); p7/p1 cleavage site (Q to R (CAG to CGG) at P3', A to V (GCT to CTT) at P2').               | Otto95, Winslow95          |
| L 10 F            | CTC to TTC   | Protease Inhibitor | DMP 450         | Y        | ?       |                   |                      | KNI-272: 7-fold; L10F/I47V/I84V: 19-fold.   | Yoshimura99                |
| L 10 F            | CTC to TTC   | Protease Inhibitor | JE-2147         | Y        | ?       |                   |                      | Ritonavir: 9-fold L10F/M46I/I47V/I84V: 28-fold, >50 passages required for isolation of resistant virus.   |                            |
| L 10 F            | CTC to TTC   | Protease Inhibitor | SC-55389A       | Y        | ?       | 2.8               |                      | N88S/L10F: 25-fold  | Potts94, Pillay96, Smidt97 |
| L 10 F            | CTC to TTC   | Protease Inhibitor | VB 11,328       | Y        | ?       |                   |                      | L10F/I84V: 8-fold   | Paraledis95                |
| L 10 F            | CTC to TTC   | Protease Inhibitor | VX-478 (141W94) | Y        | ?       |                   |                      |   | Tisdale96                  |

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold resistance | Cross-resist (-fold)  | Comments   | Refs               |
|-------------------|--------------|--------------------|-------------------------------|----------|---------|------------------|---|--|--------------------|
| L 10 F            | CTC to TTC   | Protease Inhibitor | XM323                         |          |         |                  | L10F/ V82A: 2-fold; L10F/ K45I/ I84V: 50-fold   |  | King95             |
| L 10 I            | CTC to ATC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                  |   |  | Condra96           |
| L 10 I            | CTC to ATC   | Protease Inhibitor | Ro 31-8959 (saquinavir)       | Y        |         |                  | Found in combination with G48V in vivo.   |  | Schapiro96         |
| L 10 R            | CTC to CGC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | N        | Y       |                  | XM-323 (15) L10R/ M46I/ L63P/ V82T: 4-fold; L10R/ M46I/ L63P/ V82T/ I84V: 8-fold  |  | Condra96, Condra95 |
| L 10 V            | CTC to GTC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                  | A-80987 (4)   |  | Condra96, Condra95 |
| G 16 E            | GGG to GAG   | Protease Inhibitor | ABT-378                       | Y        | ?       |                  | Passage 17 virus: Passage 17 virus: 184V/ L10F/ M46I/ ritonavir: 21-fold; T91S/ V32I/ I47V/ V47A/ G16E/ saquinavir: 4-fold H69Y: 338 fold (in presence of p7/p1 (AN/F to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation). |  | Carillo98          |
| K 20 M            | AAG to ATG   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                  | VX-478 (8)  |  | Condra96           |
| K 20 M            | AAG to ATG   | Protease Inhibitor | Nelfinavir                    | ?        | Y       |                  |   | Seen in two patients following a switch from saquinavir. Associated with reduced susceptibility to both saquinavir and nelfinavir. | Lawrence99         |
| K 20 R            | AAG to AGG   | Protease Inhibitor | ABT-538 (ritonavir)           | N        | Y       |                  |   | K20R/M36I/I54V/V82A: 41-fold   | Molla96            |
| K 20 R            | AAG to AGG   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                  | Ro-31-8959 (8);   |  | Condra96           |
| L 23 I            | CTA to ATA   | Protease Inhibitor | BILA 2185 BS                  | Y        | ?       |                  | Ro-31-8959 (50); L10F/ L23I/ V32I/ M46I/ I47V/ I54M/ L-735,524 (80); A71V/ I84V: 1,500-fold. Associated BILA 1906 BS  |  | Croteau97, Doyon96 |
| L 24 I            | TTA to ATA   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                  | Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'; p7/p1 (360) P3' to P3'', A to V (GCT to CTT) at P2').  |  | Condra96, Condra95 |
| L 24 V            | TTA to GTA   | Protease Inhibitor | SC-52151                      | Y        | ?       | 10-20            | SC-55389A   | L24V/ G48V/ A71V/ V75I/ P81T: 1000-fold  | Potts94, Pillay96  |
|                   |              |                    |                               |          |         |                  | Reviews   |  |                    |

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound              | In vitro | In vivo | -Fold -resistance (fold) | Cross-resist Comments  | Refs                 |
|-------------------|--------------|--------------------|-----------------------|----------|---------|--------------------------|--|----------------------|
| D 30 N            | GAT to AAT   | Protease Inhibitor | AG1343 (nefnavir)     | Y        | Y       |                          | D30N/A1IV: 7-fold; D30N and N88D are most common in vivo after 24 weeks of therapy; they do not cause cross-resistance to other protease inhibitors  | Patick96, Patick97   |
| V 32 I            | GTA to ATA   | Protease Inhibitor | A-77003               | Y        | ?       | 7 (enzyme resist.)       | V32I appears first; progression to V32I/M46V and V32I/ M46V/ A71V/ V82A occurs even in the absence of drug   | Kaplan94             |
| V 32 I            | GTA to ATA   | Protease Inhibitor | ABT-378               | Y        | ?       |                          | Passage 17 virus: I84V/ L10F/ M46L/ T91S/ V32I/ V32I/ 147V: Carrillo98   |                      |
|                   |              |                    |                       |          |         |                          | ritonavir, 21-fold; 46 fold. Passage 17 virus: I84V/ L10F/ saquinavir, 4-fold M46L/ T91S/ V32I/ 147V/ V47A/ G16E/ H69Y; 338 fold (in presence of p7/p1 (AN/F to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation). |                      |
| V 32 I            | GTA to ATA   | Protease Inhibitor | ABT-538 (ritonavir)   | Y        | ?       | 40                       | V32I and V82I are synergistic mutations yielding 20-fold enzyme resistance   | Molla96              |
| V 32 I            | GTA to ATA   | Protease Inhibitor | BILA 1906 BS          | Y        | ?       |                          | V32I/ A71V: 3-fold; V32I/ M46I/L/ A71V/ I84V: 5-fold; V32I/ M46I/L/ A71V/ I84A: 520-fold. 32I/ 46L/ 71V/ 84A are functionally impaired. Associated C $\alpha$ g mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'))                  | Lamarre94, Croteau97 |
| V 32 I            | GTA to ATA   | Protease Inhibitor | BILA 2011 (palinavir) | Y        | ?       | 1200                     | BILA 1906 (1400)<br>Other mutations found in p1/ p6 cleavage site  | Lamarre95            |

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound                       | In vitro | In vivo | -Fold -resistance  | Cross-resist (-fold)  | Comments   | Refs      |
|-------------------|--------------|--------------------|--------------------------------|----------|---------|--------------------|---|--|-----------|
| V 32 I            | GTA to ATA   | Protease Inhibitor | BILA 2185 BS                   | Y        | ?       |                    | BILA 1906 (360)   | L10F/ L23I/ V32I/ M46I/ I47V/ I54M/ A71V/ I84V: 1,500-fold. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'); p7/p1 cleavage site (Q to R (CAG to CGG) at P3', A to V (GCT to CTT) at P2'). V32I/ M46I/ I84V: 37-fold; V32I/ L33F/ K45I/ F53L/ A71V/ I84V/ L89M: 130-fold | Croteau97 |
| V 32 I            | GTA to ATA   | Protease Inhibitor | KNI-272                        | Y        | ?       | 2                  |   | V32I/ M46I/ V82A: 3-fold; V32I/ M46I/ A71V/ V82A: 14-fold  | Condra95  |
| V 32 I            | GTA to ATA   | Protease Inhibitor | MK-639 (L-735, 524, indinavir) | Y        | Y       |                    | M36I/I54V/A71V/V82I: 8-fold; K20R/M36I/I54V/V82A: 41-fold. In vivo, V82A/F/T/S occurs first, often followed by changes at 54, 71 and 36 | Molla96  |           |
| L 33 F            | TTA to TTC   | Protease Inhibitor | ABT-538 (ritonavir)            | N        | Y       |                    | In vivo, V82 occurs first, often followed by changes at 54, 71 and 36   | Patick96   |           |
| M 36 I            | ATG to ATA   | Protease Inhibitor | ABT-538 (ritonavir)            | N        | Y       |                    | L10F/ K45I/ I84V: 50-fold   | Tisdale94  |           |
| M 36 I            | ATG to ATA   | Protease Inhibitor | AG1343 (nefnavir)              | Y        |         |                    | Seen with V82A  | Kaplan94   |           |
| K 45 I            | AAA to ATA   | Protease Inhibitor | XM323                          | Y        | ?       | 4 (enzyme resist.) | No effect on susceptibility but improves replication competency of R8Q mutant; R8K/ M46I/ G48V: 20-fold                                 | Ho94, Kaplan94   |           |
| M 46 F            | ATG to TTC   | Protease Inhibitor | A-77003                        | Y        | ?       |                    |   |  |           |
| M 46 I            | ATG to ATA   | Protease Inhibitor | A-77003                        | Y        | ?       |                    |   |  |           |

## Mutations in HIV Protease that confer drug resistance, ordered by position.

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold -resistance | Cross-resist (fold)   | Comments  | Refs       |
|-------------------|--------------|--------------------|-------------------------------|----------|---------|-------------------|---|---|------------|
| M 46 I            | to           | Protease Inhibitor | ABT-378                       | y        |         |                   |   | Passage 17 virus: I84V/ L10F/ M46I: 4 fold; I84V/ L10F/ M46I/ T91S: 12 fold; I84V/ L10F/ M46I/ T91S/ V32I/ 147V: 46 fold. Passage 17 virus: I84V/ L10F/ M46I/ T91S/ V32I/ I47V/ V47A/ G16E/ H69Y: 338 fold (in presence of p7/p1 (ANF to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation). | Carrillo98 |
| M 46 I            | ATG to ATA   | Protease Inhibitor | ABT-538 (ritonavir)           | Y        | Y       |                   |   | M46I/ L63P/ A71V/ V82F/ 184V: 27-fold   | Molla96    |
| M 46 I            | ATG to ATA   | Protease Inhibitor | AG1343 (nefnavir)             | Y        | Y       |                   |   |   | Patick96   |
| M 46 I            | ATG to ATA   | Protease Inhibitor | BILA 1906 BS                  | Y        | ?       | L 735,524 (60)    | V32I/ A71V: 3-fold; V32I/ M46I/L/ A71V/ 184V: 5-fold; V32I/ M46I/L/ A71V/ I84A: 520-fold. V32I/M46L/A71V/I84A is functionally impaired. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'))    | Croteau97, Doyon96, Lamarre94, Lamarre95  |            |
| M 46 I            | ATG to ATA   | Protease Inhibitor | BILA 2185 BS                  | Y        | ?       | BILA 1906 (360)   | L10F/ L23I/ V32I/ M46I/ 147V/ 154M/ A71V/ 184V: 1,500-fold. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1')); p7/p1 cleavage site (Q to R (CAG to CGG) at P3', A to V (GCT to CTT) at P2'). | Croteau97   |            |
| M 46 I            | ATG to ATA   | Protease Inhibitor | DMP 450                       | Y        | ?       | KNI-272: 7-fold;  | L10F/M46I/147V/184V: >50  | Otto95, Winslow95   |            |
| M 46 I            | ATG to ATA   | Protease Inhibitor | JE-2147                       | Y        | ?       | Ritonavir: 9-fold | passages required for isolation of resistant virus.   | Yoshimura99   |            |
| M 46 I            | ATG to ATA   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | N        | Y       |                   | M46I/ L63P/ V82T: 4-fold; L10R/ M46I/ L63P/ V82T: 4-fold; L10R/ M46I/ L63P/ V82T/ 184V: 8-fold  | Condra96, Condra95  |            |

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro   | In vivo | -Fold -resistance    | (-fold) | Cross-resist   | Comments | Refs                                     |
|-------------------|--------------|--------------------|-------------------------------|--|---------|----------------------|---------|--|----------|--|
| M 46 I            | ATG to ATA   | Protease Inhibitor | VB 11,328                     | Y  | ?       |                      |         | 150V/ M46I/ 147V: 20-fold  |          | Tisdale94, Partaledis95                  |
| M 46 I            | ATG to ATA   | Protease Inhibitor | VX-478 (141W94)               | Y  | ?       | Nil                  |         |  |          | Partaledis95                             |
| M 46 L            | ATG to TTG   | Protease Inhibitor | A-77003                       | Y  | ?       | 2-3 (enzyme resist.) |         |  |          | Kaplan94                                 |
| M 46 L            | ATG to TTG   | Protease Inhibitor | BILA 1906 BS                  | Y  | ?       |                      |         |  |          | Croteau97, Doyon96, Lamarre94, Lamarre95 |
| M 46 L            | ATG to TTG   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | Y  | Y       |                      |         |  |          | Tisdale94                                |
| M 46 L            | ATG to CTG   | Protease Inhibitor | XM323                         | Y  | ?       |                      |         |  |          |  |
| M 46 V            | ATG to GTG   | Protease Inhibitor | A-77003                       | Y  | ?       |                      |         | V82A/ M46L: 7-fold; V82A/ M46L/ L97V: 11-fold  | King95   |  |
| I 47 V            | ATA to GTA   | Protease Inhibitor | ABT-378                       | Y  | ?       |                      |         | V32I appears first; progression to V32I/M46V and V32I/ M46V/A71V/V82A occurs even in the absence of drug.  |          | Tisdale94                                |
| I 47 V            | ATA to CTA   | Protease Inhibitor | BILA 2185 BS                  | Y  | ?       |                      |         | Passage 17 virus: I84V/ L10F/ M46I/ T91S/ V32I/ I47V: ritonavir, 21-fold; 46 fold, Passage 17 virus: I84V/ L10F/ saquinavir, 4-fold M46I/ T91S/ V32I/ I47V/ V47A/ G16E/ H69Y: 338 fold (in presence of p7/p1 (AN/F to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation). |          | Carrillo98                               |
| I 47 V            | ATA to CTA   | Protease Inhibitor | BILA 1906 (360)               | L10F/ L23I/ V32I/ M46I/ I47V/ I54M/ A71V/ I84V: 1,500-fold. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'); p7/p1 cleavage site (Q to R (CAG to CGG) at P3', A to V (GCT to CTT) at P2'). |         |                      |         |  |          | Croteau97                                |

## Mutations in HIV Protease that confer drug resistance, ordered by position.

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold resistance | Cross-resist (-fold) | Comments  | Refs                 |
|-------------------|--------------|--------------------|-------------------------------|----------|---------|------------------|----------------------|---|----------------------|
| I 54 M            | ATT to ATG   | Protease Inhibitor | BILA 2185 BS                  | Y        | ?       |                  |                      | BILA 1906 (360) L10F/L23I/V32I/M46I/L47V/L54M/A71V/I84V: 1,500-fold. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'); p7/p1 cleavage site (Q to R (CAG to CGG) at P3'; A to V (GCT to CTT) at P2'). | Croteau97            |
| I 54 V            | ATC to GTC   | Protease Inhibitor | ABT-538 (ritonavir)           | N        | Y       |                  |                      | I54V/V82T: 9-fold; K20R/M36I/I54V/V82A: 41-fold; M36I/I54V/A71V/V82T: 8-fold; I54V/A71V/V82A/L90N: 7-fold; In vivo, V82A/F/T/S occurs first, followed by changes at 54, 71 and 36   | Molla96              |
| I 54 V            | ATC to GTC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                  |                      |   | Lamarre94            |
| I 54 V            | ATA to GTA   | Protease Inhibitor | Ro 31-8959 (saquinavir)       | Y        |         |                  |                      | In subtype O  | Jacobsen94, Eberle95 |
| I 54 V            | ATC to GTC   | Protease Inhibitor | Ro 31-8959 (saquinavir)       | Y        |         |                  |                      | In subtype B  | Jacobsen94, Eberle95 |
| K 55 R            | AAA to AGA   | Protease Inhibitor | Nelfinavir                    | ?        | Y       |                  |                      | Seen in one patient following a switch from saquinavir. Associated with reduced susceptibility to both saquinavir and nelfinavir.   | Lawrence99           |
| R 57 K            | AGA to AAA   | Protease Inhibitor | Nelfinavir                    | ?        | Y       |                  |                      | Seen in one patient following a switch from saquinavir. Associated with reduced susceptibility to both saquinavir and nelfinavir.   | Lawrence99           |
| D 60 E            | GAT to GAA   | Protease Inhibitor | DMP 450                       | Y        | ?       |                  |                      | Probably compensatory   | Ott95, Winslow95     |
| L 63 P            | CTC to CCC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | N        | Y       |                  |                      | M46I/L63P/V82T: 4-fold; L10R/M46I/L63P/V82T/I84V: 8-fold; L10R/M46I/L63P/V82T: 4-fold   | Condra96, Condra95   |
| L 63 P            | CTC to CCC   | Protease Inhibitor | Nelfinavir                    | ?        | Y       |                  |                      | D30N/M36I/L63P: 60-fold   | Pattick98            |

## Mutations in HIV Protease that confer drug resistance, ordered by position.

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold -resistance | Cross-resist (-fold) | Comments  | Refs                                     |
|-------------------|--------------|--------------------|-------------------------------|----------|---------|-------------------|----------------------|---|--|
| H 69 Y            | CAT to TAT   | Protease Inhibitor | ABT-378                       | Y        | ?       |                   |                      | Passage 17 virus: 184V/ L10F/ M46L/ ritonavir, 21-fold; T91S/ V32I/ I47V/ V47A/ G16E/ saquinavir, 4-fold H69Y: 338 fold (in presence of p7/p1 (AN/F to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation).                                   | Carillo98                                |
| A 71 T            | GCT to ACT   | Protease Inhibitor | BMS 186,318                   | Y        | ?       |                   |                      |   | Patrick95, Rose94                        |
| A 71 T            | GCT to ACT   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                   |                      |   | Condra96, Condra95                       |
| A 71 V            | GCT to GTT   | Protease Inhibitor | A-77003                       | Y        | ?       |                   |                      | V32I appears first; progression to V32I/M46V and V32I/M46V/ A71V/V82A occurs even in the absence of drug; M46I/ L63P/ A71V/ V82F/ I84V: 27-fold   | Tisdale94, King95                        |
| A 71 V            | GCT to GTT   | Protease Inhibitor | ABT-538 (ritonavir)           | Y        | Y       |                   |                      |   | Molla96                                  |
| A 71 V            | GCT to GTT   | Protease Inhibitor | AG1343 (nefnavir)             | Y        | ?       | 5                 |                      | D30N/ A71V: 7-fold; M46I/ L63P/ A71V/ 184V: 30-fold   | Patrick98                                |
| A 71 V            | GCT to GTT   | Protease Inhibitor | BILA 1906 BS                  | Y        | ?       |                   |                      | V32I/ A71V: 3-fold; V32I/ M46I/L/ A71V/ 184V: 5-fold; V32I/ M46I/L/ A71V/ 184A: 520-fold. 32I/ 46L/ 71V/ 84A are functionally impaired. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'))  | Croteau97, Doyon96, Lamarre94, Lamarre95 |
| A 71 V            | GCT to GTT   | Protease Inhibitor | BILA 2011 (palinavir)         | Y        | ?       |                   |                      | BILA 2185: 30-fold  | Lamarre94                                |
| A 71 V            | GCT to GTT   | Protease Inhibitor | BILA 2185 BS                  | Y        | ?       |                   |                      | BILA 1906 (360) L10F/ L23I/ V32I/ M46I/ 147V/ 154M/ A71V/ 184V: 1,500-fold. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1')); p7/p1 cleavage site (Q to R (CAG to CGG) at P3'; A to V (GCT to CTT) at P2'). V32I/ M46L/ A71V/ V82A: 14-fold | Croteau97                                |
| A 71 V            | GCT to GTT   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | Y        | Y       |                   |                      |   | Tisdale94                                |

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold -resistance (-fold)  | Cross-resist Comments  | Refs              |
|-------------------|--------------|--------------------|-------------------------------|----------|---------|--|--|-------------------|
| A 71 V            | GCT to GTT   | Protease Inhibitor | SC-52151                      | Y        | ?       | Not L-735,524  | A71V/ V75I/ P81T: 20- to 30-fold; L24V/ G48V/ A71V// V75I/ P81T: 1000-fold; N88D or H11V/ M46L/ F53L/ A71V/ N88D: 10- to 20-fold | Potts94, Pillay96 |
| G 73 S            | GGT to GCT   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       | Emerges following a switch from saquinavir to indinavir.   | Duijoust97   |                   |
| G 73 S            | GGT to AGT   | Protease Inhibitor | Nelfinavir                    | ?        | Y       | Seen in two patients following a switch from saquinavir. Associated with reduced susceptibility to both saquinavir and nelfinavir. | Lawrence99   |                   |
| V 75 I            | GTA to ATA   | Protease Inhibitor | SC-52151                      | Y        | ?       | L24V/ G48V/ A71V// V75I/ P81T: 1000-fold; A71V/ V75I/ P81T: 20- to 30-fold; L24V/ G48V/ A71V// V75I/ P81T: 1000-fold               | Potts94, Pillay96  |                   |
| V 77 I            | GTA to ATA   | Protease Inhibitor | AG1343 (nelfinavir)           | Y        | Y       |  | Patrick98  |                   |
| P 81 T            | CCT to ACT   | Protease Inhibitor | SC-52151                      | Y        | ?       | A71V/ V75I/ P81T: 20- to 30-fold; L24V/ G48V/ A71V// V75I/ P81T: 1000-fold   | Potts94, Pillay96  |                   |
| I 82 T            | ATC to ACC   | Protease Inhibitor | A-77003                       | Y        | ?       | G48V/ I82T: 100-fold 82T was derived from in vitro passage of 82I)   | Swanson94  |                   |
| V 82 A            | GTC to GCC   | Protease Inhibitor | A-77003                       | Y        | ?       | Rare; seen with M46F; V32I appears first; progression to V32I/ M46V and V32I/ M46V/ A71V/ V82A occurs even in the absence of drug  | Tisdale94, Borman95, Swanson94   |                   |
| V 82 A            | GTC to GCC   | Protease Inhibitor | ABT-538 (ritonavir)           | N        | Y       | In vivo, V82 occurs first, often followed by changes at I54, A71 and M36   | Molla96  |                   |
| V 82 A            | GTC to GCC   | Protease Inhibitor | BMS 186,318                   | Y        | ?       | A71T/ V82A: 15-fold  | Patrick95, Rose94  |                   |
| V 82 A            | GTC to GCC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | Y        | Y       | V32I/ M46L/ V82A; 3-fold; V32I/ M46L/ A71V/ V82A: 14-fold  | Condra96, Condra95   |                   |
| V 82 A            | GTC to GCC   | Protease Inhibitor | Nelfinavir                    | ?        | Y       |  | Lawrence99   |                   |
| V 82 A            | GTC to GCC   | Protease Inhibitor | P9941                         | Y        | ?       | 6-8  | Otto93   |                   |

## Mutations in HIV Protease that confer drug resistance, ordered by position.

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold -resistance | Cross-resist (-fold) | Comments  | Refs                             |
|-------------------|--------------|--------------------|-------------------------------|----------|---------|-------------------|----------------------|---|----------------------------------|
| V 82 A            | GTC to GCC   | Protease Inhibitor | Ro 31-8959 (saquinavir)       | ?        | Y       |                   |                      | Follows G48V during saquinavir therapy or after a switch to nefnavir or indinavir.  | Winters97, Eastman97, Schapiro97 |
| V 82 A            | GTC to GCC   | Protease Inhibitor | SC-52151                      | Y        | ?       |                   |                      |   | Potts94, Pillay96                |
| V 82 A            | GTC to GCC   | Protease Inhibitor | SKF108922                     | Y        | ?       |                   |                      |   | Shao95                           |
| V 82 A            | GTC to GCC   | Protease Inhibitor | XM323                         | Y        | ?       |                   |                      |   | King95                           |
| V 82 F            | GTC to TTC   | Protease Inhibitor | ABT-538 (ritonavir)           | Y        | Y       |                   |                      |   |                                  |
| V 82 F            | GTC to TTC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | ?        | Y       |                   |                      |   | Paraledis94                      |
| V 82 F            | GTC to TTC   | Protease Inhibitor | XM323                         | Y        | ?       |                   |                      |   | King95                           |
| V 82 I            | GTC to ATC   | Protease Inhibitor | A-77003                       | Y        | ?       |                   |                      | No resistance alone but V32I and V82I are synergistic mutations yielding 20-fold enzyme resistance 82T was derived from in vitro passage of 82I | Kaplan94                         |
| V 82 I            | GTC to ATC   | Protease Inhibitor | XM323                         | Y        | ?       | < 2               |                      |   | King95                           |
| V 82 S            | GTC to TCC   | Protease Inhibitor | ABT-538 (ritonavir)           | N        | Y       | 6                 |                      | In vivo, V82 occurs first, often followed by changes at I54, A71 and M36  | Molla96                          |
| V 82 T            | GTC to ACC   | Protease Inhibitor | ABT-538 (ritonavir)           | N        | Y       | 3                 |                      | In vivo, V82 occurs first, often followed by changes at I54, A71 and M36; V82T has reduced replication efficacy in natural background           | Molla96                          |
| V 82 T            | GTC to ACC   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | N        | Y       |                   |                      | M46I/ L63P/ V82T: 4-fold; L10R/ M46I/ L63P/ V82T: 4-fold; L10R/ M46I/ L63P/ V82T/ I84V: 8-fold  | Condra96, Condra95               |
| V 82 T            | GTC to ACC   | Protease Inhibitor | SKF108922                     | Y        | ?       |                   |                      |   | Shao95                           |
| V 82 T            | GTC to ACC   | Protease Inhibitor | SKF108922                     | Y        | ?       |                   |                      |   | Shao95                           |

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound              | In vitro | In vivo | -Fold -resistance   | Cross-resist (-fold)   | Comments   | Refs                                     |
|-------------------|--------------|--------------------|-----------------------|----------|---------|---|--|--|--|
| I 84 A            | ATA to GCA   | Protease Inhibitor | BILA 1906 BS          | Y        | ?       | (200)   | BILA 2185 BS   | V32I/ A71V: 3-fold; V32I/ M46I.L/ A71V/ 184V: 5-fold; V32I/ M46I.L/ A71V/ 184A: 520-fold. 32I/ 46L/ 71V/ 84A are functionally impaired. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1')) | Croteau97, Doyon96, Lamarre94, Lamarre95 |
| I 84 A            | ATG to ATA   | Protease Inhibitor | BILA 2011 (palinavir) | Y        | ?       | (400); Ro 31-8959   | I84A is the most common mutation   |  | Lamarre94                                |
| I 84 V            | ATA to GTA   | Protease Inhibitor | ABT-378               | Y        | ?       | Passage 17 virus: 184V/ L10F/ M46I: 4 fold, I84V/ L10F/ ritonavir, 21-fold; M46I/ T91S/ V32I/ 147V: 46 fold, Pas-saquinavir, 4-fold | M46I/ T91S: 12 fold, I84V/ L10F/ T91S/ V32I/ 147V: 46 fold, Passage 17 virus: 184V/ L10F/ M46I/ T91S/ V32I/ 147V/ V47A/ G16E/ H69Y: 338 fold (in presence of p7/p1 (AN/F to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation). | Carillo96  |  |
| I 84 V            | ATA to GTT   | Protease Inhibitor | ABT-538 (ritonavir)   | Y        | Y       | M46I/ L63P/ A71V/ V82F/ 184V: 27-fold; V82F/ 184V: 8- to 10-fold; M46I/ L63P/ A71V/ V82F/ 184V: 27-fold                             | M46I/ L63P/ A71V/ V82F/ 184V: 30-fold  | Molla96  |  |
| I 84 V            | ATA to GTT   | Protease Inhibitor | AG1343 (nefnavir)     | ?        |         | M46I/ L63P/ A71V/ V82F/ 184V: 30-fold   |  | Patick96   |  |
| I 84 V            | ATA to GTT   | Protease Inhibitor | BILA 1906 BS          | Y        | ?       | BILA 2185 BS(200)   | V32I/ A71V: 3-fold; V32I/ M46I.L/ A71V/ 184V: 5-fold; V32I/ M46I.L/ A71V/ 184A: 520-fold. 32I/ 46L/ 71V/ 84A are functionally impaired. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'))                                       | Croteau97, Doyon96, Lamarre94, Lamarre95   |  |

## Mutations in HIV Protease that confer drug resistance, ordered by position.

| Amino Acid | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold -resistance | Cross-resist (fold)  | Comments   | Refs  |          |
|------------|--------------|--------------------|-------------------------------|----------|---------|-------------------|--|--|---|----------|
| I 84 V     | ATA to GTA   | Protease Inhibitor | BILA 2185 BS                  | Y        | ?       |                   | BILA 1906 BS(360)  | L10F/ L23V/ V32I/ M46I/ I47V/ I54M/ A71V/ I84V: 1,500-fold. Associated Gag mutations: p1/p6 cleavage site (L to F (CTT to TTT at P1'); p7/p1 cleavage site (Q to R (CAG to CGG) at P3', A to V (GCT to CTT) at P2'). Minor resistance mutation for BMS-232632. | Croteau97   |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | BMS-232632                    | Y        | ?       |                   |  |  | Otto95, Winslow95   |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | DMP 450                       | Y        | ?       |                   | KNI-272: 7-fold; Ritonavir: 9-fold                                 | L10F/I47V/I84V: 19-fold. L10F/M46I/I47V/I84V: 28-fold. >50 passages required for isolation of resistant virus.   | Yoshimura99   |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | JE-2147                       | Y        | ?       |                   | G48V/ I84V/ L90M: 30-fold; L10R/ M46L/ L63P/ V82T/ I84V: 8-fold    |  | Condra96, Condra95  |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | MK-639 (L-735,524, indinavir) | N        | Y       |                   |  |  |   |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | MP-134                        | Y        | ?       | 10                | MP-167(5) ABT-538(10) MK-639(8) SC-5215(8) Ro31-895(2) (IC90 data) |  | Mo96  |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | Ro 31-8959 (saquinavir)       | Y        | ?       |                   |  |  | Tisdale94   |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | RPI-312                       | Y        | ?       | 5                 |  |  | el-Farrash94  |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | SKFI108842                    | Y        | ?       |                   |  |  | Shao95  |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | VB 11,328                     | Y        | ?       |                   |  |  | Partaledis95  |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | VX-478 (141W94)               | Y        | ?       |                   |  |  | Partaledis95  |          |
| I 84 V     | ATA to GTA   | Protease Inhibitor | XM323                         | Y        | ?       | 12                | P9941; not A-77003 or Ro 31-8959                                   | V82F/ I84V: 92-fold; L10F/ K45I/ I84V: 50-fold   | Tisdale94, King95   |          |
| N 88 D     | AAT to GAT   | Protease Inhibitor | AG1343 (nelfinavir)           | Y        | Y       |                   |  |  | D30N and N88D are most common in vivo after 24 weeks of therapy; they do not cause cross-resistance to other protease inhibitors. | Patick96 |

**Mutations in HIV Protease that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug      | Compound                      | In vitro | In vivo | -Fold -resistance | (-fold)                     | Cross-resist  | Comments         | Refs              |
|-------------------|--------------|--------------------|-------------------------------|----------|---------|-------------------|-----------------------------|---|------------------|-------------------|
| N 88 D            | AAT to GAT   | Protease Inhibitor | SC-52151                      | Y        | ?       |                   |                             | N88D compensatory, no resistance alone  |                  | Potts94, Pillay96 |
| N 88 S            | AAT to AGT   | Protease Inhibitor | BMS-232632                    | Y        | ?       |                   |                             | Major resistance mutation for BMS-232632.   |                  | Gong99            |
| N 88 S            | AAT to AGT   | Protease Inhibitor | SC-55389A                     | Y        | ?       | 20                | L735,S724 (3); not SC-52151 | N88S/L10F: 25   |                  | Smidt97           |
| L 90 M            | TTG to ATG   | Protease Inhibitor | ABT-538 (ritonavir)           | N        | Y       |                   |                             | 82A/ 54V/ I/ 71V/ 90L/ M: 7-fold  |                  | Molla96           |
| L 90 M            | TTG to ATG   | Protease Inhibitor | AG1343 (nelfinavir)           | N        | Y       |                   |                             |   | Rare in patients |                   |
| L 90 M            | TTG to ATG   | Protease Inhibitor | MK-639 (L-735,S24, indinavir) | ?        | Y       |                   |                             |   |                  | Condra96          |
| L 90 M            | TTG to ATG   | Protease Inhibitor | Ro 31-8959 (saquinavir)       | Y        | Y       |                   |                             | G48V/ L90M: >100-fold enzyme resistance; double mutant rare in vivo; L90M most common in vivo; G48V/ I84V/ L90M: 30-fold  |                  | Jacobsen94        |
| T 91 S            | ACT to TCT   | Protease Inhibitor | ABT-378                       | Y        | ?       |                   |                             | Passage 17 virus: I84V/ L10F/ M46I/ T91S: 12 fold, ritonavir, 21-fold; I84V/ L10F/ M46I/ T91S/ V32I/ I47V: saquinavir, 4-fold 46 fold, Passage 17 virus: I84V/ L10F/ M46I/ T91S/ V32I/ I47V/ V47A/ G16E/ H69Y: 338 fold (in presence of p7/p1 (AN/F to VN/F) cleavage-site mutation and p1/p6 (F/L to F/F) cleavage-site mutation). |                  | Carrillo98        |
| L 97 V            | TTA to GTA   | Protease Inhibitor | XM323                         | Y        | ?       |                   |                             | No resistance alone; V82A/ L97V: 3-fold; V82A/ M46I/ L97V: 11-fold  |                  | King95            |

**Mutations in HIV Integrase that confer drug resistance, ordered by position.**

| Amino Acid | Codon      | Class of Drug       | Compound        | In vitro | In vivo | -Fold -resistance (-fold) | Cross-resist   | Comments | Ref's  |
|------------|------------|---------------------|-----------------|----------|---------|---------------------------|--|----------|--------|
| Change     | Change     |                     |                 |          |         |                           |  |          |        |
| G 140 S    | GGC to AGC | Integrase inhibitor | L-Chicoric Acid | Y        | ?       | 156-fold                  | Mutation located in the catalytic core of integrase. Mildly attenuates virus growth. |          | King98 |

**Mutations in HIV Env that confer drug resistance, ordered by position.**

| Amino Acid Change | Codon Change | Class of Drug            | Compound              | In vitro | In vivo | -Fold resistance | Cross-resist (-fold) | Comments | Refs   |
|-------------------|--------------|--------------------------|-----------------------|----------|---------|------------------|----------------------|----------|--|
| R 22 A            | AGG to AGA   | Fusion/Binding Inhibitor | RPR103611             | Y        | ?       |                  |                      |          | Labrosse97   |
| G 36 S            | GGT to AGT   | Fusion/Binding Inhibitor | DP178 (T20)           | Y        | ?       |                  |                      |          | Rimsky98   |
| V 38 M            | GTG to ATG   | Fusion/Binding Inhibitor | DP178 (T20)           | Y        | ?       |                  |                      |          | Rimsky98   |
| I 84 S            | ATC to AGC   | Fusion/Binding Inhibitor | RPR103611             | Y        | ?       |                  |                      |          | Labrosse97   |
| N 106 K           | AAT to AAG   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                  |                      |          | Schols98   |
| S 113 N           | AGT to AAT   | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?       |                  |                      |          | AMB2763: 3-fold.   |
| S 134 N           | AGC to AAC   | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?       |                  |                      |          | S113N/ S134N/ K269E/ Q278E/ N293D/ N323S/ R387I: 250-fold; 11.3 is in the V1 loop region |
| S 134 N           | AGC to AAC   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                  |                      |          | V2 loop region; S113N/ S134N/ K269E/ Q278E/ N293D/ N323S/ R387I: 250-fold                |
| F 145 L           | TTC to TTA   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       |                  |                      |          | SDF-1 $\beta$ : 15-fold; 106K/ 134N/ 145L/ 245I/ 269E/ 278H/ AMB2763: 3-fold.            |
| F 145 L           | TTC to TTA   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                  |                      |          | 288V/ 293D/ 364-367 Deletion/ 387T: 15-fold.   |
| N 188 K           | AAT to AAA   | Fusion/Binding Inhibitor | Siamycin I            | Y        | ?       |                  |                      |          | N188K/ G332E/ N351D/ A550T/ N633D/ L762S: 9-fold   |
| I 228 V           | ATA to GTA   | Fusion/Binding Inhibitor | JM-2763               | Y        | ?       |                  |                      |          | Combination of mutations: 2- to 100-fold   |
| F 245 I           | TTC to ATC   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                  |                      |          | DeVreese96, DeVreese96a  |
| F 245 I           | TTC to ATC   | Fusion/Binding Inhibitor |                       |          |         |                  |                      |          | AMB2763: 3-fold.   |
| K 269 E           | AAA to GAA   | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?       |                  |                      |          | V3 loop region; S113N/ S134N/ K269E/ Q278E/ N293D/ N323S/ R387I: 250-fold                |

## Mutations in HIV Env that confer drug resistance, ordered by position.

| Amino Acid Change | Codon Change | Class of Drug            | Compound              | In vitro | In vivo | -Fold resistance  | Cross-resist (-fold)  | Comments | Refs                    |
|-------------------|--------------|--------------------------|-----------------------|----------|---------|-------------------|---|----------|-------------------------|
| N 269 E           | AAC to GAA   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                   | SDF-1 $\beta$ : 15-fold; 106K/ 134N/ 145L/ 245L/ 269E/ 278H/ AMB2763: 3-fold. |          | Schols98                |
| N 270 S           | AAT to AGT   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       |                   | 288V/ 293D/ 364-367 Deletion/ 387T; 15-fold.                                  |          | DeVreese96, DeVreese96a |
| R 272 T           | AGA to ACA   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       |                   |   |          | DeVreese96, DeVreese96a |
| S 274 R           | AGT to AGA   | Fusion/Binding Inhibitor | JM-2763               | Y        | ?       |                   |   |          | DeVreese96, DeVreese96a |
| S 274 R           | AGT to AGA   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       | DS (> 7 to 6,667) | Combination of mutations: 95- to 792-fold                                     |          | DeVreese96, DeVreese96a |
| Q 278 H           | CAG to CAT   | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?       |                   | V3 loop region: S113N/ S134N/ K269E/ Q278E/ N293D/ N323S/ R387I: 250-fold     |          | Este97, Este96a         |
| Q 278 H           | CAG to CAT   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       |                   |   |          | DeVreese96, DeVreese96a |
| Q 278 H           | CAG to CAC   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                   | SDF-1 $\beta$ : 15-fold; 106K/ 134N/ 145L/ 245L/ 269E/ 278H/ AMB2763: 3-fold. |          | DeVreese96, DeVreese96a |
| Q 278 H           | CAG to CAT   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       |                   | 288V/ 293D/ 364-367 Deletion/ 387T; 15-fold.                                  |          | Schols98                |
| I 288 V           | ATA to GTA   | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?       |                   |   |          | DeVreese96, DeVreese96a |
| I 288 V           | ATA to GTC   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                   | SDF-1 $\beta$ : 15-fold; 106K/ 134N/ 145L/ 245L/ 269E/ 278H/ AMB2763: 3-fold. |          | DeVreese96, DeVreese96a |
| N 293 D           | AAT to GAT   | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?       |                   | 288V/ 293D/ 364-367 Deletion/ 387T; 15-fold.                                  |          | Este97, Este96a         |
| N 293 D           | AAT to GAT   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?       |                   |   |          | Este97, Este96a         |
| N 293 H           | AAT to CAT   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       |                   | V3 loop region: S113N/ S134N/ K269E/ Q278E/ N293D/ N323S/ R387I: 250-fold     |          | DeVreese96, DeVreese96a |
| A 297 T           | GCA to ACA   | Fusion/Binding Inhibitor | JM-2763               | Y        | ?       |                   | SDF-1 $\beta$ : 15-fold; 106K/ 134N/ 145L/ 245L/ 269E/ 278H/ AMB2763: 3-fold. |          | DeVreese96, DeVreese96a |
| A 297 T           | GCA to ACA   | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?       |                   | 288V/ 293D/ 364-367 Deletion/ 387T; 15-fold.                                  |          | Este97, Este96a         |

## Mutations in HIV Env that confer drug resistance, ordered by position.

| Amino Acid | Codon      | Change                   | Class of Drug         | Compound | In vitro | In vivo | -Fold -resistance | (fold) | Cross-resist  | Comments                | Refs |
|------------|------------|--------------------------|-----------------------|----------|----------|---------|-------------------|--------|---|-------------------------|------|
| N 323 S    | AAT to AGT | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?        |         |                   |        | C3 region; S113N/ S134N/ K269E/ Q278E/ N293D/ N323S/ R387I: 250-fold          | Este97, Este96a         |      |
| G 332 E    | GGA to GAA | Fusion/Binding Inhibitor | Siamycin I            | Y        | ?        |         |                   |        | N188K/ G332E/ N351D/ A550T/ N633D/ L762S: 9-fold                              | Lin96                   |      |
| N 351 D    | AAT to GAT | Fusion/Binding Inhibitor | Siamycin I            | Y        | ?        |         |                   |        | N188K/ G332E/ N351D/ A550T/ N633D/ L762S: 9-fold                              | Lin96                   |      |
| P 385 L    | CCA to CTA | Fusion/Binding Inhibitor | JM-2763               | Y        | ?        |         |                   |        |   | DeVreese96, DeVreese96a |      |
| P 385 L    | CCA to CTA | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?        |         |                   |        |   | DeVreese96, DeVreese96a |      |
| R 387 I    | AGA to ACA | Fusion/Binding Inhibitor | Dextran sulphate (DS) | Y        | ?        |         |                   |        | CD4 binding region; S113N/ S134N/ K269E/ Q278E/ N293D/ N323S/ R387I: 250-fold | Este97, Este96a         |      |
| R 387 T    | AGA to ACA | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?        |         |                   |        | SDF-1 $\beta$ : 15-fold; 106K/ 134N/ 145L/ 245L/ 269E/ 278H/ AMB2763: 3-fold. | Schols98                |      |
| Q 410 E    | CAA to GAA | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?        |         |                   |        |   | DeVreese96, DeVreese96a |      |
| S 433 P    | TCC to CCC | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?        |         |                   |        |   | DeVreese96, DeVreese96a |      |
| V 457 I    | GTA to ATA | Fusion/Binding Inhibitor | JM-3100 (SID791)      | Y        | ?        |         |                   |        |   | DeVreese96, DeVreese96a |      |
| A 550 T    | GCC to ACC | Fusion/Binding Inhibitor | Siamycin I            | Y        | ?        |         |                   |        | N188K/ G332E/ N351D/ A550T/ N633D/ L762S: 9-fold                              | Lin96                   |      |
| N 633 D    | AAT to GAT | Fusion/Binding Inhibitor | Siamycin I            | Y        | ?        |         |                   |        | N188K/ G332E/ N351D/ A550T/ N633D/ L762S: 9-fold                              | Lin96                   |      |
| L 762 S    | TTG to TCG | Fusion/Binding Inhibitor | Siamycin I            | Y        | ?        |         |                   |        | N188K/ G332E/ N351D/ A550T/ N633D/ L762S: 9-fold                              | Lin96                   |      |
| 364-368    | Deletion   | Fusion/Binding Inhibitor | SDF-1 $\alpha$        | Y        | ?        |         |                   |        | SDF-1 $\beta$ : 15-fold; 106K/ 134N/ 145L/ 245L/ 269E/ 278H/ AMB2763: 3-fold. | Schols98                |      |
|            |            |                          |                       |          |          |         |                   |        | 288V/ 293D/ 364-367 Deletion/ 387T: 15-fold.                                  |                         |      |

**Mutations in SIVRT that confer drug resistance, ordered by position.**

| Amino Acid | Codon      | Class of Drug               | Compound | In vitro | In vivo | -Fold -resistance | Cross-resist (-fold)  | Comments  | Refs   |
|------------|------------|-----------------------------|----------|----------|---------|-------------------|---|---|--|
| Change     | Change     | Change                      | Change   |          |         |                   |   |   |  |
| K 65 R     | AAA to AGA | SIV Nucleoside RT Inhibitor | PMPA     | ?        | Y       | 5                 | 3TC (80); ddC; d4T; PMEA and 1118V. Observed changes at N69S and 1118V do not result in increased resistance. | K65R appears first, followed by N69S and 1118V. Observed changes at N69S and 1118V do not result in increased resistance. | VanRompay'96, Cherrington'96a, VanRompay'97a |
| Q 151 M    | CAG to ATG | SIV Nucleoside RT Inhibitor | AZT      | ?        | Y       | >100              | ddI; ddC; d4T; 3TC  |   | VanRompay'97                                 |
| M 184 V    | ATG to GTG | SIV Nucleoside RT Inhibitor | (-)FTC   | Y        | ?       |                   |   |   | Schinazi'95                                  |

**Mutations in FIVRT that confer drug resistance, ordered by position.**

| Amino Acid | Codon      | Class of Drug               | Compound | In vitro | In vivo | -Fold -resistance | Cross-resist (-fold)      | Comments   | Refs              |
|------------|------------|-----------------------------|----------|----------|---------|-------------------|---------------------------|--|-------------------|
| Change     | Change     | Change                      | Change   |          |         |                   |                           |  |                   |
| D 3 H      | GAT to CAT | FIV Nucleoside RT Inhibitor | ddC      | Y        | ?       | 4                 | ddI; PFA                  |  | Medlin'96, Zhu'96 |
| V 47 I     | GTA to ATA | FIV Nucleoside RT Inhibitor | d4T      | Y        | ?       | 4-6               | PFA (>50); AZT; ddI; PMEA |  | Smith'96          |
| P 156 S    | CCA to TCA | FIV Nucleoside RT Inhibitor | 3TC      | Y        | ?       | 7                 | AZT (4), AZT + 3TC (6)    |  | Smith'98          |
| M 183 T    | ATG to ACG | FIV Nucleoside RT Inhibitor | (-)FTC   | Y        | ?       | 10                | ddC                       | Corresponds to 184 in HIV; M183V recombinant displays 10-fold resistance to 3TC or (-)FTC. | Smith'97          |

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**Abbreviations**

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**Amino acids**

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|   |               |
|---|---------------|
| A | alanine       |
| C | cysteine      |
| D | aspartate     |
| E | glutamate     |
| F | phenylalanine |
| G | glycine       |
| H | histidine     |
| I | isoleucine    |
| K | lysine        |
| L | leucine       |
| M | methionine    |
| N | asparagine    |
| P | proline       |
| Q | glutamine     |
| R | arginine      |
| S | serine        |
| T | threonine     |
| V | valine        |
| W | tryptophan    |
| Y | tyrosine      |

**Compounds**

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|                                 |  |
|---------------------------------|--|
| 1592U89                         | (1 <i>S</i> ,4 <i>R</i> )-4-[2-amino-6-cyclopropyl-amino]-9 <i>H</i> -purin-9-yl]-2-cyclopentene-1-methanol succinate (a carbovir analogue, Glaxo Wellcome)  |
| 3TC                             | (-)- $\beta$ -L-2',3'-dideoxy-3'-thiacytidine (Glaxo Wellcome)   |
| 1737                            | Tetrahydronaphthalene lignan derivative  |
| $\alpha$ -APA R18893            | $\alpha$ -nitro-anilino-phenylacetamide  |
| A-77003, A-75925<br>and A-80987 | C2 symmetry-based protease inhibitors (Abbott Laboratories)  |
| AAP-BHAP                        | bisheteroarylpirperazine analogue (Pharmacia & Upjohn)   |
| ABT-378                         | Protease inhibitor   |
| ABT-538                         | C2 symmetry-based protease inhibitor (Abbott Laboratories)   |
| ADAMII                          | Methyl 3',3''-dichloro-4',4''-dimethoxy-5',5''-bis(methoxycarbonyl)-6,6-diphenyl-5-hexenoate. (An alkenyldiarylmethane).   |
| AZdU                            | 3'-azido-2',3'-dideoxyuridine  |
| AZT                             | 3'-azido-3'-deoxythymidine (Glaxo Wellcome)  |
| AZT-p-ddI                       | 3'-azido-3'-deoxythymidilyl-(5',5')-2',3'-dideoxyinosinic acid (Ivax)  |
| BHAP                            | bisheteroarylpirperazine   |
| BILA 1906                       | <i>N</i> -{1 <i>S</i> -[[[3-[2 <i>S</i> -(1,1-dimethylethyl)amino]carbonyl-4 <i>R</i> -]3-pyridinylmethyl]thio]-1-piperidinyl]-2 <i>R</i> -hydroxy-1 <i>S</i> -(phenylmethyl)propyl]amino]carbonyl]-2-methylpropyl}-2-quinolinecarboxamide (Bio-Mega/Boehringer Ingelheim) |

**Abbreviations (cont)****Compounds (cont)**

|                  |  |
|------------------|--|
| BILA 2185        | <i>N</i> -(1-dimethylethyl)-1-[2 <i>S</i> -[[2-2,6-dimethoxyphenoxy)-1-oxoethyl]amino]-2 <i>R</i> -hydroxy-4-phenylbutyl]4 <i>R</i> -pyridinylthio)-2-piperidine-carboxamide (Bio-Mega/Boehringer Ingelheim)     |
| BM+51.0836       | thiazolo-isoindolinone derivative  |
| BMS 186,318      | aminodiol derivative HIV-1 protease inhibitor (Bristol-Myers Squibb)   |
| BMS-232632       | An azapeptide protease inhibitor   |
| d4API            | 9-[2,5-dihydro-5-(phosphonomethoxy)-2-furanyl]adenine (Gilead Sciences)  |
| d4C              | 2',3'-didehydro-2',3'-dideoxycytidine  |
| d4T              | 2',3'-didehydro-3'-deoxythymidine (Bristol-Myers Squibb)   |
| ddC              | 2',3'-dideoxycytidine (Roche)  |
| ddI              | 2',3'-dideoxyinosine (Bristol-Myers Squibb)  |
| DMP 266          | a 1,4-dihydro-2 <i>H</i> -3,1-benzoxazin-2-one   |
| DMP 450          | [4 <i>R</i> -(4- $\alpha$ ,5- $\alpha$ ,6- $\beta$ ,7- $\beta$ )]-hexahydro-5,6-bis(hydroxy)-1,3-bis(3-amino)phenyl)methyl)-4,7-bis(phenylmethyl)-2 <i>H</i> -1,3-diazepin-2-one-bismesylate (Avid Therapeutics) |
| DP178            | Synthetic peptide containing amino acids 127–162 of HIV-1 gp41   |
| DXG              | (-)- $\beta$ -D-dioxolane-guanosine  |
| EBU-dM           | 5-ethyl-1-ethoxymethyl-6-(3,5-dimethylbenzyl)uracil  |
| E-EBU            | 5-ethyl-1-ethoxymethyl-6-benzyluracil  |
| DS               | dextran sulphate   |
| E-EPSeU          | 1-(ethoxymethyl)-(6-phenylselenyl)-5-ethyluracil   |
| E-EPU            | 1-(ethoxymethyl)-(6-phenyl-thio)-5-ethyluracil   |
| F-ddA            | 2'-fluoro-2',3'-dideoxyadenosine   |
| (-)-FTC          | (-)- $\beta$ -L-2',3'-dideoxy-5-fluoro-3'-thiacytidine (Triangle Pharmaceuticals)  |
| GW420867X        | S-3-ethyl-6-fluoro-4-isopropoxycarbonyl-3,4-dihydro-quinoxalin-2(1 <i>H</i> )-one  |
| HBY 097          | ( <i>S</i> )-4-isopropoxycarbonyl-6-methoxy-3-(methylthio-methyl)-3,4-dihydroquinoxalin-2(1 <i>H</i> )-thione  |
| HEPT             | 1-[(2-hydroxyethoxy)methyl]6-(phenylthio)thymine   |
| JE-2147          | An allophenylnorstatine-containing dipeptide protease inhibitor  |
| JM2763           | 1,1'-(1,3-propanediyl)-bis-1,4,8,11-tetraazacyclo-tetradecane (Johnson Matthey)  |
| JM3100           | 1,1'-[1,4-phenylenebis-(methylene)]bis-(1,4,8,11-tetraazacyclotetradecane) octahydrochloride dihydrate (Johnson Matthey)   |
| KNI-272          | (2 <i>S</i> ,3 <i>S</i> )-3-amino-2-hydroxy-4-phenylbutyric acid-containing tripeptide   |
| L-697,593        | 5-ethyl-6-methyl-3-(2-phthalimido-ethyl)pyridin-2(1 <i>H</i> )-one   |
| L-697,661        | 3-[(4,7-dichloro-1,3-benzoxazol-2-yl)methyl]amino-5-ethyl-6-methylpyridin-2(1 <i>H</i> )-one   |
| L-chichoric acid | Integrase inhibitor  |
| L-FDDC           | (-)- $\beta$ -L-5-fluoro-2',3'-dideoxy-cytidine  |
| L-FDOC           | (-)- $\beta$ -L-5-fluoro-dioxolane cytosine  |
| MK-639           | hydroxy-aminopentane amide HIV-1 protease inhibitor (Merck & Co)   |
| MKC442           | 6-benzyl-1-ethoxymethyl-5-isopropyluracil (I-EBU, Triangle Pharmaceuticals/Mitsubishi)   |
| MP-134           | C2 symmetry-based protease inhibitor   |
| MP-167           | C2 symmetry-based protease inhibitor   |

**Abbreviations (cont)****Compounds (cont)**

|                       |  |
|-----------------------|--|
| nevirapine            | 11-cyclopropyl-5,11-dihydro-4-methyl-6 <i>H</i> -dipyridol[3,2-b:2',3'-e]diazepin-6-one (Boehringer Ingelheim)   |
| NNRTI                 | non-nucleoside reverse transcriptase inhibitor   |
| NSC648400             | 1-benzoyloxymethyl-5-ethyl-6-(alpha-pyridylthio)uracil (E-BPTU)  |
| P9941                 | [2-pyridylacetyl-IlePheAla-y(CHOH)] <sub>2</sub> (Dupont Merck)  |
| PFA                   | phosphonoformate (foscarnet, Astra)  |
| PMEA                  | 9-(2 phosphonylmethoxyethyl)adenine (Gilead Sciences)  |
| PMPA                  | ( <i>R</i> )-9-(2-phosphonyl-methoxypropyl)adenine (Gilead Sciences)   |
| QM96521               | 1,1,3-trioxo-2 <i>H</i> ,4 <i>H</i> -thieno[2,4-3][1,2,4]thiadiazine derivative (TTD)  |
| Ro 31-8959            | hydroxyethylamine derivative HIV-1 protease inhibitor (Roche)  |
| RPI-312               | 1-[(3 <i>S</i> )-3-(n-alpha-benzoyloxycarbonyl)-l-aspargyl]-amino-2-hydroxy-4-phenyl-butryl]- <i>n</i> -tert-butyl-l-proline amide (peptidyl protease inhibitor) |
| RPR103611             |  |
| RT                    | reverse transcriptase  |
| S-1153                | 5-(3,5-dichlorophenyl)thio-4-isopropyl-1-(4-pyridyl)methyl-1 <i>H</i> -imidazol-2-yl methyl carbamate  |
| S-2720                | 6-chloro-3,3-dimethyl-4-(isopropenyl-oxycarbonyl)-3,4-dihydro-quinoxalin-2(1 <i>H</i> )thione  |
| SC-52151              | hydroxyethylurea isostere protease inhibitor (Searle)  |
| SC-55389A             | hydroxyethyl-urea isostere protease inhibitor (Searle)   |
| SDF-1 $\alpha$        | Stromal cell-derived factor 1 $\alpha$   |
| TIBO R82150           | (+)-(5 <i>S</i> )-4,5,6,7-tetrahydro-5-methyl-6-(3-methyl-2-butenyl)-imidazo[4,5,1- <i>jk</i> ][1,4]-benzodiazepin-2(1 <i>H</i> )-thione (Janssen)               |
| TIBO 82913            | (+)-(5 <i>S</i> )-4,5,6,7-tetrahydro-9-chloro-5-methyl-6-(3-methyl-2-butenyl)-imidazo-[4,5,1- <i>jk</i> ]-[1,4]benzo-diazepin-2(1 <i>H</i> )-thione (Janssen)    |
| TSAO-m <sup>3</sup> T | [2',5'-bis- <i>O</i> -(tert-butyl-dimethylsilyl)-3'-spiro-5'-(4'-amino-1',2'-oxathiole-2',2'-dioxide)]- $\beta$ -D-pentofuranosyl-N <sup>3</sup> -methylthymine  |
| U-90152               | 1-[3-[(1-methylethyl)-amino]-2-pyridinyl]-4-[[5-[(methylsulphonyl)-amino]-1 <i>H</i> -indol-2yl]carbonyl]-piperazine   |
| U-95133               | (Alkylamino)piperidine bis(heteroaryl) piperazine analog   |
| U-104489              | (Alkylamino)piperidine bis(heteroaryl) piperazine analog   |
| UC-040                | thiocarboxanilide derivative (Uniroyal Chemical Co)  |
| UC                    | thiocarboxanilide derivatives (Uniroyal Chemical Co)   |
| UC-781                | <i>N</i> -[4-chloro-3-(3-methyl-2-butenyloxy)phenyl]-2-methyl-3-furan-carbothioamide   |
| UC-82                 | <i>N</i> -[4-chloro-3-(3-methyl-2-butenyloxy)phenyl]-2-methyl-3-thiophene-carbothioamide   |
| VB 11,328             | hydroxyethyl-sulphonamide protease inhibitor (Vertex Pharmaceuticals)  |
| VX-478                | hydroxyethylsulphonamide protease inhibitor (Vertex Pharmaceuticals)   |
| XM 323                | cyclic urea protease inhibitor (Dupont Merck)  |

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