... moves to amend H.F. No. 2987 as follows:

Page 1, line 10, delete "4" and insert "3"

Page 2, line 5, after "4" insert ", or a prescription drug that can only be dispensed to a patient registered with the drug's manufacturer in accordance with federal Food and Drug Administration requirements"

Page 2, line 14, after "drugs" insert "and medical supplies"

Page 2, after line 17, insert:

"(i) "Original, sealed, unopened, tamper-evident packaging" means packaging that is sealed, unopened, and tamper-evident, including a manufacturer's original unit dose or unit-of-use container, a repackager's original unit dose or unit-of-use container, or unit-dose packaging prepared by a licensed pharmacy according to the standards of Minnesota Rules, part 6800.3750."

Page 2, line 18, delete "(i)" and insert "(j)"

Page 3, line 28, delete "prescription drug" and insert "health" and after "drug" insert "or medical supply"

Page 3, line 30, after "drugs" insert "or medical supplies"

Page 4, line 27, delete "drug's lot number and"

Page 5, line 28, delete "lot number;"

Page 5, line 30, delete ", lot number;"

Page 6, line 6, after "drugs" insert "or supplies"

Page 6, line 19, after "substances" insert "or prescription drugs that can only be dispensed to a patient registered with the drug's manufacturer"
Page 6, line 25, delete "recalled" and after "supply" insert "that is the subject of a Class I or Class II recall"

Page 6, line 27, after the period insert "A drug that potentially is subject to a recall need not be destroyed if its packaging bears a lot number and that lot of the drug is not subject to the recall. If no lot number is on the drug's packaging, it must be destroyed."

Page 6, line 33, after "destroyed:" insert "and"

Page 7, line 1, delete "; and" and insert a period

Page 7, delete line 2

Page 7, line 20, after "drug" insert "or supply"