- WAC 182-530-7000 Reimbursement. (1) The agency's reimbursement for a prescription drug dispensed through point-of-sale (POS) must not exceed the lesser of actual acquisition cost (AAC) plus a professional dispensing fee or the provider's usual and customary charge.
- (2) The agency selects the sources for pricing information used to set POS AAC.
 - (3) The POS AAC is calculated as the lowest of:
 - (a) National average drug acquisition cost (NADAC);
 - (b) Maximum allowable cost (MAC);
 - (c) Federal upper limit (FUL);
- (d) 340B Actual acquisition cost (340B AAC) for drugs purchased under section 340B of the Public Health Service (PHS) Act (see WAC 182-530-7900 for exceptions); or
 - (e) Automated maximum allowable cost (AMAC).
- (4) Where NADAC does not exist, other available reference prices from national sources such as wholesale acquisition cost, or average manufacturer price will be used as the basis of the reimbursement.
- (5) Where NADAC does not accurately reflect the actual acquisition costs in Washington state, a percentage adjustment to NADAC will be made to the reimbursement.
- (6) The agency may set POS AAC for specified drugs or drug categories at a maximum allowable cost other than that determined in subsection (2) of this section based on specific product acquisition costs. The agency considers product acquisition costs in setting a rate for a drug or a class of drugs.
- (7) The agency bases POS AAC drug reimbursement on the actual package size dispensed.
- (8) The agency reimburses a pharmacy for the least costly dosage form of a drug within the same route of administration, unless the prescriber has designated a medically necessary specific dosage form or the agency has selected the more expensive dosage form as a preferred drug.
- (9) If the pharmacy provider offers a discount, rebate, promotion or other incentive which directly relates to the reduction of the price of a prescription to the individual nonmedicaid customer, the provider must similarly reduce its charge to the agency for the prescription.
- (10) If the pharmacy provider gives an otherwise covered product for free to the general public, the pharmacy must not submit a claim to the agency.
 - (11) The agency does not reimburse for:
- (a) Prescriptions written on presigned prescription blanks filled out by nursing facility operators or pharmacists;
 - (b) Prescriptions without the date of the original order;
- (c) Drugs used to replace those taken from a nursing facility emergency kit;
 - (d) Drugs used to replace a physician's stock supply;
- (e) Outpatient drugs, biological products, insulin, supplies, appliances, and equipment included in other reimbursement methods including, but not limited to:
 - (i) Diagnosis-related group (DRG);
 - (ii) Ratio of costs-to-charges (RCC);
 - (iii) Nursing facility daily rates;
 - (iv) Managed care capitation rates;
 - (v) Block grants; or

- (vi) Drugs prescribed for clients who are on the agency's hospice program when the drugs are related to the client's terminal illness and related condition.
- (f) Hemophilia and von Willebrand related products shipped to clients for administration in the home unless the products are provided through a qualified hemophilia treatment center of excellence (COE) as defined in WAC 182-531-1625.

[Statutory Authority: RCW 41.05.021 and 41.05.160. WSR 17-07-001, § 182-530-7000, filed 3/1/17, effective 4/1/17. Statutory Authority: RCW 41.05.021. WSR 12-16-061, § 182-530-7000, filed 7/30/12, effective 11/1/12. WSR 11-14-075, recodified as § 182-530-7000, filed 6/30/11, effective 7/1/11. Statutory Authority: RCW 74.04.050, 74.08.090, 74.09.530, and 74.09.700. WSR 07-20-049, § 388-530-7000, filed 9/26/07, effective 11/1/07.]