

A bill for an act

relating to human services; changing the pharmacy dispensing fee; amending  
Minnesota Statutes 2006, section 256B.0625, subdivision 13e.

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF MINNESOTA:

Section 1. Minnesota Statutes 2006, section 256B.0625, subdivision 13e, is amended to  
read:

Subd. 13e. **Payment rates.** (a) The basis for determining the amount of payment  
shall be the lower of the actual acquisition costs of the drugs plus a fixed dispensing  
fee; the maximum allowable cost set by the federal government or by the commissioner  
plus the fixed dispensing fee; or the usual and customary price charged to the public.  
The amount of payment basis must be reduced to reflect all discount amounts applied  
to the charge by any provider/insurer agreement or contract for submitted charges to  
medical assistance programs. The net submitted charge may not be greater than the patient  
liability for the service. Effective July 1, 2007, the pharmacy dispensing fee shall be  
\$3.65 for single-source drugs and \$12.92 for multiple-source generic drugs, except that  
the dispensing fee for intravenous solutions which must be compounded by the pharmacist  
shall be \$8 per bag, \$14 per bag for cancer chemotherapy products, and \$30 per bag  
for total parenteral nutritional products dispensed in one liter quantities, or \$44 per bag  
for total parenteral nutritional products dispensed in quantities greater than one liter.  
An inflation adjustment shall be made annually to the dispensing fee for multiple-source  
generic prescriptions based on the CPI-all items for urban consumers. Actual acquisition  
cost includes quantity and other special discounts except time and cash discounts. The  
actual acquisition cost of a drug shall be estimated by the commissioner, at average  
wholesale price minus 12 percent. The actual acquisition cost of antihemophilic factor

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drugs shall be estimated at the average wholesale price minus 30 percent. The maximum allowable cost of a multisource generic drug may be set by the commissioner and it shall be comparable to, but no ~~higher than, the maximum amount paid by other third-party payors in this state who have maximum allowable cost programs~~ lower than the price of the drug available to retail pharmacies for purchase from prescription drug wholesalers. Establishment of the amount of payment for drugs shall not be subject to the requirements of the Administrative Procedure Act.

(b) Effective for fee-for-service pharmacy services rendered on or after July 1, 2007, the commissioner may, within the limits of available appropriation, increase dispensing fees to pharmacists and pharmacies deemed by the commissioner to be critical access pharmacy providers. Reimbursement to a critical access pharmacy provider may be increased by not more than 50 percent above the dispensing fee that would otherwise be paid to the provider. In determining which pharmacists and pharmacies shall be deemed critical access pharmacy providers, the commissioner shall review:

(1) the utilization rate in the service area in which the pharmacist or pharmacy operates for pharmacy services to patients covered by medical assistance, general assistance medical care, or MinnesotaCare as their primary source of coverage;

(2) the level of services provided by the pharmacist or pharmacy to patients covered by medical assistance, general assistance medical care, or MinnesotaCare as their primary source of coverage; and

(3) whether the level of services provided by the pharmacist or pharmacy is critical to maintaining adequate levels of patient access within the service area.

~~(b)~~ (c) An additional dispensing fee of \$.30 may be added to the dispensing fee paid to pharmacists for legend drug prescriptions dispensed to residents of long-term care facilities when a unit dose blister card system, approved by the department, is used. Under this type of dispensing system, the pharmacist must dispense a 30-day supply of drug. The National Drug Code (NDC) from the drug container used to fill the blister card must be identified on the claim to the department. The unit dose blister card containing the drug must meet the packaging standards set forth in Minnesota Rules, part 6800.2700, that govern the return of unused drugs to the pharmacy for reuse. The pharmacy provider will be required to credit the department for the actual acquisition cost of all unused drugs that are eligible for reuse. Over-the-counter medications must be dispensed in the manufacturer's unopened package. The commissioner may permit the drug clozapine to be dispensed in a quantity that is less than a 30-day supply.

3.1 ~~(c)~~ (d) Whenever a generically equivalent product is available, payment shall be on  
3.2 the basis of the ~~actual acquisition cost of~~ federal upper limit set for the generic drug, or on  
3.3 the maximum allowable cost established by the commissioner.

3.4 ~~(d)~~ (e) The basis for determining the amount of payment for drugs administered in  
3.5 an outpatient setting shall be the lower of the usual and customary cost submitted by the  
3.6 provider or the amount established for Medicare by the United States Department of  
3.7 Health and Human Services pursuant to title XVIII, section 1847a of the federal Social  
3.8 Security Act.

3.9 ~~(e)~~ (f) The commissioner may negotiate lower reimbursement rates for specialty  
3.10 pharmacy products than the rates specified in paragraph (a). The commissioner may  
3.11 require individuals enrolled in the health care programs administered by the department  
3.12 to obtain specialty pharmacy products from providers with whom the commissioner has  
3.13 negotiated lower reimbursement rates. Specialty pharmacy products are defined as those  
3.14 used by a small number of recipients or recipients with complex and chronic diseases  
3.15 that require expensive and challenging drug regimens. Examples of these conditions  
3.16 include, but are not limited to: multiple sclerosis, HIV/AIDS, transplantation, hepatitis  
3.17 C, growth hormone deficiency, Crohn's Disease, rheumatoid arthritis, and certain forms  
3.18 of cancer. Specialty pharmaceutical products include injectable and infusion therapies,  
3.19 biotechnology drugs, high-cost therapies, and therapies that require complex care. The  
3.20 commissioner shall consult with the formulary committee to develop a list of specialty  
3.21 pharmacy products subject to this paragraph. In consulting with the formulary committee  
3.22 in developing this list, the commissioner shall take into consideration the population  
3.23 served by specialty pharmacy products, the current delivery system and standard of care in  
3.24 the state, and access to care issues. The commissioner shall have the discretion to adjust  
3.25 the reimbursement rate to prevent access to care issues.