
THIRD READING

Bill No: SB 958
Author: Limón (D) and Portantino (D)
Amended: 4/18/22
Vote: 21

SENATE HEALTH COMMITTEE: 8-0, 4/6/22
AYES: Pan, Eggman, Hurtado, Leyva, Limón, Roth, Rubio, Wiener
NO VOTE RECORDED: Melendez, Gonzalez, Grove

SENATE APPROPRIATIONS COMMITTEE: 5-2, 5/19/22
AYES: Portantino, Bradford, Kamlager, Laird, Wieckowski
NOES: Bates, Jones

SUBJECT: Medication and Patient Safety Act of 2022

SOURCE: California Hospital Association

DIGEST: This bill restricts the ability of health plans/insurers, or their designated medical groups or pharmacy benefit managers, from requiring or incentivizing patients to have infused or injected medications supplied by a vendor to the patient, or to the patient's physician office, clinic, infusion center, or hospital outpatient department, rather than maintained at the location where the infused or injected medication will be administered.

ANALYSIS:

Existing law:

- 1) Establishes the Department of Managed Health Care (DMHC) to regulate health plans under the Knox-Keene Health Care Service Plan Act of 1975 (Knox-Keene Act) and the California Department of Insurance (CDI) to regulate health insurers. [HSC §1340, et seq., and INS §106, et seq.]

- 2) Establishes the California Department of Public Health (CDPH) and sets forth its powers and duties, including, but not limited to, licensing and regulating health care facilities, including acute care hospitals. [HSC §1250, et seq.]
- 3) Prohibits a health plan contract from requiring or allowing a health care service provider (but not other providers such as health facilities, hospices, or surgical centers) to assume or be at any financial risk for any specified covered injectable medications and adult vaccines that are administered in the office of a physician and surgeon or prescribed by a physician and surgeon for self-administration by the patient. Requires these items to be reimbursed on a fee-for-service basis at the negotiated contract rate or through an alternate funding mechanism mutually agreed to by the health plan and the health care service provider, subject to any applicable copayment or deductible, by the health plan. [HSC §1375.8]
- 4) Permits a health care service provider to assume financial risk for the items described above after making the request in writing at the time of negotiating an initial contract or renewing a contract with a health plan. Prohibits a health plan from requesting or requiring as a condition of the contract agreement a health care service provider to assume the financial risk for any of those items. [HSC §1375.8]

This bill:

- 1) Prohibits a health plan/insurer or its designee (such as a medical group, pharmacy benefit manager, or other designee from requiring a vendor (pharmacy) to dispense an infused or injected medication directly to a patient with the intent that the patient will transport the medication to a health care provider for administration. Indicates a vendor is not an integrated health system's internal pharmacy that dispenses a patient's prescription medical and transports the product to the health system's location of drug administration.
- 2) Defines an infused or injected medication as an outpatient prescription medication or biologic, other than a vaccine, that cannot reasonably be self-administered by the patient or other individual assisting the patient, and is typically administered by a licensed health care provider acting within the scope of their professional licensure in a physician's office, clinic, infusion center, or hospital outpatient department.

- 3) Permits a health plan/health insurer or its designee to arrange for an infused or injected medication to be administered to an enrollee/insured in the enrollee's/insured's home when the treating health care provider and patient determines administration in the home setting is in the best interest of the patient and requires the provider to document it in the patient's medical record.
- 4) Prohibits a health plan/health insurer or its designee from requiring, as a condition of coverage or payment, or offering an incentive for, an infused or injected medication to be supplied by a vendor specified by the plan/insurer or designee unless all of specified requirements are true.
- 5) Requires a health plan/insurer or designee that implements a policy requiring, as a condition of coverage or payment, that an infused or injected medication be supplied by a specified vendor to provide written notification to the treating health care provider, the entity authorized to contract for the provider's services, and the facility in which the provider administers the medication, at least 45 business days in advance of the effective date of the proposed change.
- 6) Permits the treating health care provider, the entity authorized to contract for the provider's services, and the facility in which the provider administers the medication the right to negotiate this material change to the contract. Requires, if the parties to the contract cannot agree, any party to have the right to terminate the contract before the implementation of the change.
- 7) Prohibits a health plan/insurer or its designee from interfering with the enrollee's/insured's right to obtain a covered, medically necessary infused or injected medication from a participating provider of the enrollee's/insured's choosing. Prohibits a health plan/insurer or its designee from refusing to authorize or approve, exclude coverage for, deny payment for, or offer an incentive for, an infused or injected medication administered by a participating provider based on the site of service, whether the site is a physician's office, clinic, infusion center, or hospital outpatient department.

Comments

According to the author, this bill ensures that patients with complex and life-threatening illnesses who need specialty, infused or injected medications administered by a doctor or nurse, such as chemotherapy, can receive the necessary medication in a safe, timely manner. This bill does not prohibit the current practice of requiring hospitals and physicians to order specialty medications from a third-

party pharmacy, but rather sets appropriate guardrails to ensure patient safety and medication integrity.

Drug trends. According to IQVIA’s 2023 outlook report the global pharmaceutical market will exceed \$1.5 trillion by 2023 growing at a 3–6% compound annual growth rate over the next five years. The key drivers of growth will continue to be the U.S. and pharmerging markets with 4–7% and 5–8% compound annual growth, respectively. In the U.S., overall spending growth is driven by a range of factors including new product uptake and brand pricing, while it is offset by patent expiries and generics. New products and losses of exclusivity will continue to drive similar dynamics across developed markets, while product mix will continue to shift to specialty and orphan products. An average of 54 new active substance launches per year are expected over the next five years and two-thirds of launches will be specialty products, lifting specialty share of spending to near 50% by 2023 in most developed markets. At the same time, the impact of losses of exclusivity in developed markets is expected to be \$121 billion between 2019 and 2023, with 80% of this impact, or \$95 billion, in the U.S. By 2023, biosimilar competition in the biologics market will be nearly three-times larger than it is today. This will result in approximately \$160 billion in lower spending over the next five years than it would have if biosimilars did not enter the market.

Background on white bagging. The National Association of Pharmacy Boards (NAPB) describes “white bagging” as the distribution of patient-specific medications from a pharmacy, typically a specialty pharmacy, to the physician’s office, hospital, or clinic for administration. It is often used in oncology practices to obtain costly injectable medications that are distributed by specialty pharmacies and may not be available in all non-specialty pharmacies. “Brown bagging” is the dispensing of medication from a pharmacy (typically a specialty pharmacy) directly to a patient, who then transports the medications to the physician’s office for administration. In a 2018 report on these topics, the NABP indicates that as the specialty pharmacy model becomes more prevalent more patient care will be subject to white and brown bagging under mandates by third party payers. The NAPB concluded that there is a legitimate patient protection issue when a specialty drug is distributed to an entity other than the patient.

FISCAL EFFECT: Appropriation: No Fiscal Com.: Yes Local: Yes

According to the Senate Appropriations Committee:

DMHC. The DMHC estimates the total cost of this bill to be approximately \$619,000 Managed Care Fund (MCF) and 2.9 PYs in FY 2022-23, \$868,000 MCF

and 3.3 PYs in FY 2023-24, \$935,000 MCF and 3.5 PYs in FY 2024-25, \$931,000 MCF and 3.5 PYs in FY 2025-26 and annually thereafter. All costs associated would be incurred by the Managed Care Fund (MCF) and covered through fees assessed on health plans.

CDI. CDI estimates \$11,000 FY 2022-23, \$20,000 FY 2023-24, \$10,000 FY 2024-25 for CDI to revise policy forms, policies and procedures, and provider contracts, and to review the revised documents when submitted.

SUPPORT: (Verified 5/17/22)

California Hospital Association (source)
Adventist Health
Alliance of Catholic Health Care
American Academy of Pediatrics
Association of California Healthcare Districts
California Children's Hospital Association
California Council for the Advancement of Pharmacy
California Life Sciences
California Pharmacists Association
California Rheumatology Alliance
California Society of Health System Pharmacists
Cedars-Sinai
Centinela Hospital Medical Center
Children Now
Children's Specialty Care Coalition
City of Hope National Medical Center
Coast Plaza Hospital
Cottage Health
Dignity Health
District Hospital Leadership Forum
Encompass Health Rehabilitation Hospital of Bakersfield
Henry Mayo Newhall Hospital
Huntington Hospital
Infusion Providers Alliance
John C Fremont Healthcare District
John Muir Health
Kaweah Delta Healthcare District
Lucile Packard Children's Hospital
Marshall Medical Center
MemorialCare Health System

PIH Health
Rady Children's Hospital San Diego
Sacramento Behavioral Healthcare Hospital
Santa Rosa Behavioral Healthcare Hospital
Scripps Health
Sierra View Medical Center
Stanford Health Care
Sutter Health
Tahoe Forest Health System
Tenet Health
Tenet Healthcare Corporation
United Hospital Association
University of California
US Oncology Network

OPPOSITION: (Verified 5/17/22)

America's Health Insurance Plans
Association of California Life and Health Insurance Companies
California Association of Health Plans
California Chamber of Commerce
Pharmaceutical Care Management Association

ARGUMENTS IN SUPPORT: This bill is sponsored by the California Hospital Association (CHA), which writes that some health plans require physicians and hospitals to order medication from a third-party vendor contracted with the health plan, on a patient-by-patient basis, instead of using medication the hospital or physician's offices have in stock. As such, these policies threaten patient and medication safety since they are not part of a hospital's or physician's office's quality control protocols, the provider has no way of assuring drug integrity. In addition, if medications are not delivered in a timely manner or have to be re-ordered due to same-day clinical presentations, patient care can be delayed, causing them unsafe and undue stress. For more complex medications, to ensure the highest quality product, increased care and attention are crucial to ensuring product quality control is required. Specialty drugs that are compounded, or that require a clinical assessment before infusion to ensure proper strength and dosage, are especially difficult to order from a third-party vendor because what was ordered may not be what is needed at the point of care. Instead of administering available medication from their stock, providers must request them from these third-party vendors. If the vendor is unable to deliver the medication, the patient must return on a different day for the infusion. Delaying treatment may put the

patient's health at risk because many complex treatment plans involving infused medications are time sensitive, will likely lead to patient distress, and may impact the patient's overall treatment plan. CHA writes white bagging may present an initial cost savings to health plans, increased costs for hospitals and physician's offices to manage inventory on a patient-by-patient basis, plus the real patient costs of delayed care and disease progression both outweigh and potentially cancel any savings to the health care system. The California Council for the Advancement of Pharmacy writes too often, and for cost effective purposes, pharmacy benefit managers and/or health insurers require medically fragile patients to take their injectable/infused medicines to their appointment with a physician's office or outpatient hospital facility for treatment. This practice, known as white bagging, puts the patient in jeopardy because there is no quality assurance during the transport of the meds (for example, temperature sensitive drugs); however, if the same medication is already stored properly at the outpatient facility or doctor's office, then quality is not compromised for both the patient and the drug. White bagging is not necessarily a bad option for oral medications, but in the instance of injectable and infused specialty drugs, the practice of white-bagging puts the patient in a potentially health-compromised position. The Children's Specialty Care Coalition writes many patients with serious and life-threatening diseases need medications that are infused or injected at an outpatient hospital facility or in a doctor's office. Health care providers keep these specialty medications in stock, so that they are available in the form and dose whenever needed by a patient, and take primary responsibility for patient and medication safety. For more complex medications, to ensure the highest quality product, increased care and attention are crucial to ensuring product quality control is required. Specialty drugs that are compounded, or that require a clinical assessment before infusion to ensure proper strength and dosage, are especially difficult to order from a third-party vendor because what was ordered may not be what is needed at the point of care. Instead of administering available medication from their stock, providers must request them from these third-party vendors. If the vendor is unable to deliver the medication, the patient must return on a different day for the infusion. Delaying treatment may put the patient's health at risk because many complex treatment plans involving infused medications are time sensitive, will likely lead to patient distress, and may impact the patient's overall treatment plan.

ARGUMENTS IN OPPOSITION: The California Association of Health Plans, Association of California Life and Health Insurance Companies, America's Health Insurance Plans (AHIP) and PCMA (opponents) write that the drugs to which this bill applies are high-priced medications that treat complex, chronic, or rare conditions (e.g., cancer, multiple sclerosis, rheumatoid arthritis) and cannot be safely administered by a patient or their caregiver. They can have special handling

and/or administration as these include most biologic drugs. Both the number and price of specialty drugs have rapidly increased in recent years, and specialty drugs are a leading contributor to drug spending growth. The price of a specialty drug can range from thousands to ten thousands of dollars per regimen. An AHIP study found that costs per single treatment for drugs administered in hospitals were an average of \$7,000 more than those purchased through pharmacies. Hospitals, on average, charged double the prices for the same drugs, compared to pharmacies. Physician offices charged 22% higher prices for the same drugs on average. Specialty pharmacies are different from traditional “brick and mortar” pharmacies because they focus on dispensing drugs that retail pharmacies are not equipped to dispense. Specialty pharmacies typically ship products directly to clinicians just like a manufacturer or wholesaler would. They must abide by all state and federal legal and regulatory requirements, including chain of custody tracking in addition to meeting extra safety requirements for specialty drugs imposed by the FDA and drug manufactures. According to the opponents, specialty pharmacies are used only for prescription drugs that may be safely delivered this way, and they are on the hook for the cost of these very expensive drugs due to spoilage, delays, mishandling and therefore have every incentive not to waste any products. Delays can occur for individual patient deliveries in the same way delays can occur for hospitals or physician office deliveries. Health plans build additional safety features to avoid patient harm. Health plans always have exception processes in place to address circumstances of quality, safety, medical necessity, and/or care interruption. Often times medications are shipped with enough additional supply so that facilities can adjust a dose as needed at the time of administration. Because of the great cost differential between drugs provided by a specialty pharmacy and those provided by the hospital or physician’s office opponents strenuously object to this provision that allows the provider to make unilateral requests to protect their profits. Furthermore, opponents have concerns about the provision in the exception process that requires a health plan or insurer to arrange for the request to be reviewed by a pharmacist and a practicing physician in the same or similar specialty who treats the medical condition or provides the treatment for which the particular medication is used.

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