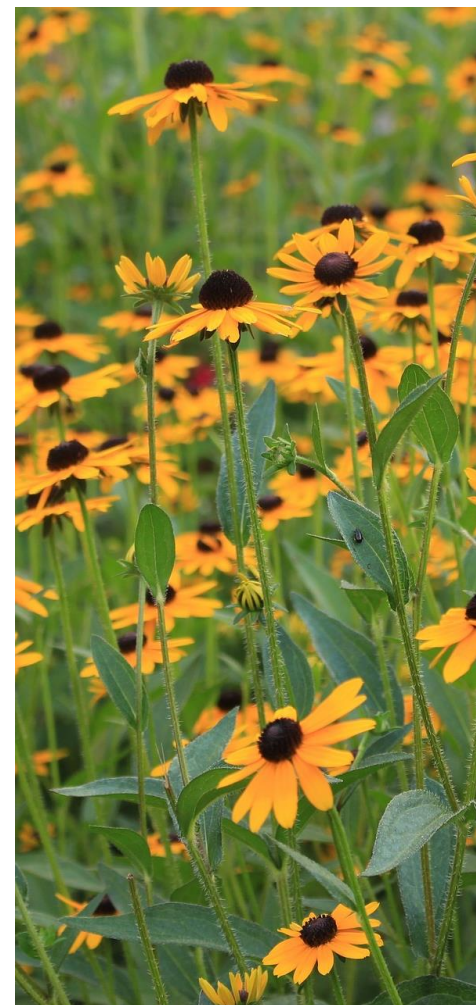




**The Hilltop Institute**

Prescription Drug  
Affordability Board (PDAB)  
Legislative Measures  
Methodology

April 26, 2022



UMBC

# Overview

- MD Code Ann., Health-Gen. § 21-2C-08
- Identify:
  - Brand name drugs with a launch Wholesale Acquisition Cost (WAC) of \$30,000+ *per year or course of treatment*
  - Brand name drugs with a WAC increase of \$3,000+ *per year or course of treatment*
  - Biosimilars with a launch WAC not at least 15% lower than the reference biologic
  - Generic drugs that cost \$100+ *for a 30-day supply (or less)* **and** experienced a price increase of at least 200%

# Challenges

- WAC pricing data are unit cost
- How to aggregate to yearly utilization?
- Scope of the data forestalls manual review
  - Tens of thousands of National Drug Codes (NDCs)
  - Recommended dosage can vary

## Proposed Methodology

- Proposed solution
  - Top-down, data-driven methodology to calculate NDC-specific utilization using claims data
- Then, link this NDC-specific utilization calculation to unit WAC pricing

# Hypothetical Example: Distribution of Claims

**Table 1. Distribution of Claims of a Hypothetical NDC in CY 2020**

<b>NDC</b>	<b>Person ID</b>	<b>Claim Date</b>	<b># Units</b>
12345878910	1	1/1/2020	1
12345878910	1	4/1/2020	1
12345878910	1	7/1/2020	1
12345878910	1	10/1/2020	1
12345878910	2	5/1/2020	1
12345878910	2	8/1/2020	1
12345878910	2	11/1/2020	1
12345878910	3	11/1/2020	1
12345878910	4	1/1/2020	1
12345878910	4	4/1/2020	1
12345878910	4	7/1/2020	1
12345878910	4	10/1/2020	1
12345878910	5	1/1/2020	1
12345878910	5	2/15/2020	1
12345878910	5	4/1/2020	1
12345878910	5	5/15/2020	1
12345878910	5	7/1/2020	1
12345878910	5	8/1/2020	1

# Hypothetical Example: Distribution of Person-Level Annual Utilization

Table 2. Distribution of Annual Utilization among Users of a Hypothetical NDC in CY 2020

NDC	Person ID	Total # Units
12345878910	1	4
12345878910	2	3
12345878910	3	1
12345878910	4	4
12345878910	5	6

Table 3: Ordered Distribution of Annual Utilization among Users of a Hypothetical NDC in CY 2020

NDC	Person ID	Total # Units
12345878910	3	1
12345878910	2	3
12345878910	1	4
12345878910	4	4
12345878910	5	6

Median annual utilization across users: **4 units**

# Estimated Utilization Distribution

- In the preceding example, use the observed median quantity (4) in conjunction with WAC prices
- Very similar methodology for generic drugs
  - Typical utilization for a 30-day supply or less

# Potential Limitations

- Restricts focus to NDCs for which there is any utilization
  - **Mitigation:** We use three large claims databases to cast as wide a net as possible
    - Maryland Medicaid
    - Maryland All-Payer Claims Database (APCD)
    - Maryland Medicare
- Relies on representativeness of claims experience
  - **Mitigation:** Manual review of a sample of drugs to assess accuracy of utilization estimate



# More Potential Limitations

- Utilization estimates may be skewed downward by individuals with partial year coverage
  - **Mitigation:** Restrict focus to individuals with at least 320 days of coverage (Medicaid and APCD).
- This analysis is NDC-level, not drug-level
  - **Mitigation:** It is not *a priori* clear how to aggregate across different dosage forms to the “drug” level. Future work could seek to reprice claims utilization using WAC prices, then aggregate to the drug level.

## Additional Assumptions

- First NDC entry in the First Databank WAC database is the launch price
- Consumer Price Index (CPI) Medical for inflation adjustment
- Identifying all generic drugs that were:
  - Ever over \$100 for a 30-day supply (or less)
  - AND**
  - Ever experienced a 200% price increase

# Biosimilars Methodology

1. Used the FDA Purple Book, current as of December 2021, to ID:
  - a. Generic name, proprietary name, and approval date of all approved biosimilars in the U.S.
  - b. Generic name and proprietary name of the corresponding reference biologic
2. Systemic search of manufacturer websites and press releases to ID biosimilar *launch dates*
3. Used the FDA directory to identify all NDC codes for *launched* biosimilars and their originator biologics

# Methodology

continued

4. Used NDC codes, launch date data, and the WAC pricing file to:
  - a. ID biosimilar *launch WAC*
  - b. ID the reference biologic WAC *at the time of biosimilar launch*
  - c. Flag biosimilars that launched at a WAC price that was not at least 15% lower than the referenced brand biologic at the time the biosimilars are launched

# Sensitivity Check

- Repeated step 4a and 4b above by biosimilar *packaging availability*.
- Example:
  - Drug A available as a 10mg/ml 50 ml vial & a 10 mg/ml 10 ml vial
  - WAC pricing is per ML → theoretically the pricing should be the same for a 10ml vial and a 50ml vial.
  - Check for potential differential pricing:
    - ID launch WAC of Drug A available in a 10ml vial formulation & in a 50ml vial formulation.
    - ID WAC/ml of reference biologic Drug B at the time of Drug A launch for the corresponding 10ml vial formulation and the corresponding 50 ml vial formulation.
    - Flag any differential pricing by packaging → **NONE.**