# ANNUAL REPORT TO THE GOVERNOR AND LEGISLATURE

# New York State Medicaid Preferred Drug Program

STATE FISCAL YEAR APRIL 1, 2015 – MARCH 31, 2016

**New York State Department of Health** 

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# **ACRONYMS**

Acronym/Term	Definition		
BLTG	Brand Less Than Generic		
CCC Clinical Call Center			
CDRP	Clinical Drug Review Program		
СРТ	Certified Pharmacy Technician		
DAW	Dispense As Written		
DOH	New York State Department of Health		
DURB	Drug Utilization Review Board		
FDA	Federal Drug Administration		
FHPlus	Family Health Plus		
FQD	Frequency, Quantity, Duration		
FUL	Federal Upper Limit		
HID Health Information Designs			
IVR	Interactive Voice Response		
МСО	Managed Care Organization		
MGDP	Mandatory Generic Drug Program		
NMPI	National Medicaid Pooling Initiative		
NYS	New York State		
P&TC Pharmacy and Therapeutics Committee			
PA	Prior Authorization		
PDL	Preferred Drug List		
PDP Preferred Drug Program			
PDSP Preferred Diabetic Supply Program			
PSL Preferred Supply List			
SDC	State Direct Contracting		
SFY	State Fiscal Year		
SMAC	State Maximum Allowable Cost		
VIPS	Voice Interactive Phone System		

### I. Background

In 2005, legislation was passed (Sections 270-277 of the Public Health Law, as enacted by Section 10 of Part C of Chapter 58 of the Laws of 2005) establishing the Medicaid Preferred Drug Program (PDP) and Clinical Drug Review Program (CDRP). The legislation expanded the membership of the Drug Utilization Review Board (DURB), established operational and administrative procedures and provided authority for the State to establish a Preferred Drug List (PDL) in order to receive supplemental rebates from drug manufacturers.

In 2006, the PDP and CDRP were implemented through a contract with Magellan Medicaid Administration (formerly known as First Health Services Corporation – FHSC). Magellan Medicaid Administration was selected through a competitive bid to operate the Clinical Call Center that supports the Medicaid PDP, CDRP, and Mandatory Generic Drug Program (MGDP); provide outreach and education services; assist with the clinical drug reviews; and obtain competitive pricing for prescription drugs through supplemental drug rebate agreements with drug manufacturers participating in the National Medicaid Pooling Initiative (NMPI). Additional programs that have been added since the inception of the Preferred Drug Program include the Brand Less Than Generic Program; Drug Utilization Program; and the Dose Optimization Program.

Effective October 1, 2008, the population eligible for the PDP was expanded to include Family Health Plus (FHPlus) members (program has since ended – 12/31/2014). The pharmacy benefit for FHPlus members was "carved-out" of the managed care plan benefit package and moved under the administration of the Medicaid fee-for-service program, whereby prescriptions for Family Health Plus members became subject to Medicaid's Preferred, Clinical Drug Review and Mandatory Generic Drug Programs and eligible for supplemental drug rebates. Effective October 1, 2011, members in mainstream Medicaid managed care and FHPlus no longer received pharmacy services through NYS Medicaid FFS Pharmacy Benefit Programs. The Department of Health (DOH) has established a goal of having virtually all Medicaid enrollees served in care management by April 2022.

Expansion of the programs and operational enhancements continued this SFY. At the end of the SFY there were a total of 108 drug classes subject to the PDP and 28 therapeutic categories warranted re-review by the DURB due to new clinical and/or financial information. Two new drug classes were reviewed for inclusion on the PDL. No new drugs were recommended by the DURB for inclusion to the CDRP.

# **II.** Program Overview

### The Role of the Drug Utilization Review Board (DURB)

The Drug Utilization Review Board (DURB) (Appendix 2), is comprised of health care professionals appointed by the Commissioner of Health and includes physicians and pharmacists that actively practice in New York. Without vacancies, the DURB consists of nineteen members, fifteen of which are clinicians, preferably with experience in at least one of the following specialties: HIV, AIDS, geriatrics, pediatrics, mental health, or internal medicine and will be comprised of the following:

- One chairperson representing the Department of Health;
- Six licensed and actively practicing physicians;
- Six licensed and actively practicing pharmacists;
- One licensed and actively practicing nurse practitioner or midwife;
- Two drug utilization review experts, at least one of who is a pharmacologist;
- Three consumers or consumer representatives of organizations with a regional or statewide constituency and who have been involved in activities related to health care consumer advocacy, including issues affecting Medicaid or EPIC recipients.

The board provides clinical guidance to the Commissioner regarding the utilization of pharmaceuticals within the Medicaid program including but not limited to, the:

- establishment and implementation of medical standards and criteria for the retrospective and prospective DUR program;
- development, selection, application, and assessment of educational interventions for physicians, pharmacists and recipients that improve care, and management of pharmacy programs including the PDP and CDRP;
- collaboration with managed care organizations to address drug utilization concerns and to implement consistent management strategies across the fee-for-service and managed care pharmacy benefits; and
- review of therapeutic classes subject to the Preferred Drug Program.

The DURB is subject to the Public Officers Law and meetings are subject to the Open Meeting Law. To ensure transparency in the process, a notice of each meeting and the agenda is posted on the DOH website thirty (30) days prior to the meeting. Interested parties are given an opportunity to submit materials to the DURB for consideration and to provide public testimony on the agenda items. The meetings are audiocast and all audiocasts are available on-demand for a minimum of 30 days.

The DURB hears public comments and first reviews clinical information relevant to the drugs under consideration during the public session. The clinical information consists of the most current therapeutic drug class reviews and evidence-based research obtained by Magellan Medicaid Administration, DOH staff and through the DOH's participation in the Oregon Health Sciences University Drug Effectiveness Review Project. Materials submitted by interested parties prior to the meeting, as well as oral testimony provided during the public session, are discussed.

Following the clinical presentation and consideration of all clinical information, the DURB may adjourn for an executive session in order to evaluate confidential drug pricing information with respect to rebates. The DUR Board reconvenes in open session to discuss any remaining issues, then votes on the recommendations to be submitted to the Commissioner of Health.

A summary of the meeting's proceedings, including the DURB's recommendations, is posted to the DOH website, which initiates a 5-day public comment opportunity. The DURB's recommendations as well as the statements made during the public comment period are then presented to the Commissioner who makes the final determination.

The Commissioner's final determination is posted to the DOH website, and includes an analysis of the impact on state public health plan populations, providers and the fiscal impact to the State.

A list of the drug classes reviewed during this fiscal year appear in Appendix 3.

### The Preferred Drug Program (PDP)

The PDP promotes utilization of clinically appropriate, cost effective prescription drugs through the use of a Preferred Drug List (PDL). Most preferred drugs on the PDL can be prescribed without any additional action taken by the prescriber; non-preferred drugs require prior authorization (PA) by calling or faxing the Clinical Call Center (CCC) or PA may also be auto assigned if clinical criteria has been met at the point of service.

PA may be required if a drug is non-preferred or to override clinical criteria including, but not limited to frequency, quantity, duration (*FQD*), diagnosis or step therapy requirements. Details regarding these limitations can be found by accessing the Preferred Drug List (PDL) at: <a href="https://newyork.fhsc.com/providers/PDP\_about.asp">https://newyork.fhsc.com/providers/PDP\_about.asp</a>

In developing the PDL, the DOH works with the DURB to select therapeutic drug classes where drugs in the class produce similar clinical effects or outcomes. The DURB evaluates the clinical effectiveness, safety and patient outcomes among drugs in the therapeutic classes chosen for review. If the DURB establishes that one drug is significantly more effective and safer than others in the class, that drug must be preferred without consideration of cost. If the DURB ascertains that there is no substantial clinical difference among the drugs in the class, it then considers the net cost of the drug after rebates as a factor in determining preferred status. The DURB also considers how its recommendations may impact current prescribing and dispensing practices and patient care. Recommendations are presented to the Commissioner of Health, who makes the final determination regarding which drugs will be listed as preferred or non-preferred.

The DOH issues the PDL (<u>Appendix 4</u>), which lists all drugs on the Preferred Drug Program. Revisions were made to the PDL to include links to other pharmacy management programs that may impact PDL drugs. The PDL is updated and posted on the website (newyork.fhsc.com) whenever there is a change.

# The Clinical Drug Review Program (CDRP)

The CDRP was implemented in October 2006 and initially applied to only three drugs: Revatio<sup>®</sup>, Serostim<sup>®</sup> and Zyvox<sup>®</sup>. The CDRP was designed to ensure specific drugs are utilized in a medically appropriate manner. The CDRP requires PA for specific drugs for which there may be

specific safety issues, public health concerns, the potential for fraud and abuse, or the potential for significant overuse and misuse.

Legislation prohibits cost as a basis for the selection of a drug for the CDRP or as a denial reason when a PA is requested.

Prior to the CDRP legislation, Serostim® and Zyvox® were subject to PA due to public health concerns and the potential for abuse through overuse and misuse. PA was obtained using an automated voice interactive phone system (VIPS). Legislation required that these drugs be transitioned to the CDRP. With that transition in October 2006, the PA process was changed from the VIPS process to the staffed clinical call center, which allows for a clinical discussion with the prescriber.

The DURB reviews drugs for inclusion to the CDRP, as needed. Their recommendations are based on review of established Food and Drug Administration (FDA) approved clinical indications, clinical research and input from interested parties. When making the final determination, the following clinical criteria are considered by the Commissioner:

- Whether the drug requires monitoring of prescribing protocols to protect both the long-term efficacy of the drug and the public health;
- The potential for, or a history of overuse, abuse, diversion or illegal utilization;
- The potential for or a history of utilization inconsistent with approved indications.

The complete list of drugs/drug classes subject to the CDRP at the end of this fiscal year is as follows:

- Anabolic Steroids
- Central Nervous System (CNS) Stimulants (for patients 18 years of age and older)
- Fentanyl Mucosal Agents
- Growth Hormone
- **Lidoderm**<sup>®</sup> (lidocaine patch 5%)
- Phosphodiesterase type-5 (PDE-5) Inhibitors for pulmonary arterial hypertension (PAH)
- Regranex® (becaplermin gel)
- Serostim<sup>®</sup> [somatropin (rDNA origin) for injection]
- Synagis<sup>®</sup> (palivizumab)
- Topical Immunomodulators
- Truvada® (emtricitabine and tenofovir disoproxil fumarate)
- Xyrem<sup>®</sup> (sodium oxybate)
- Zyvox® (linezolid) and Sivextro® (tedizolid)

# Brand Less Than Generic (BLTG) Program

On April 26, 2010, New York State Medicaid implemented a cost containment initiative, which promotes the use of certain multi-source brand name drugs when the cost of the brand name drug is less expensive than the generic equivalent. Additionally, the *Brand Less Than Generic* 

(BLTG) program is designed to promote the use of certain multi-source brand name drugs when the cost of the brand name product net of all rebates is less than its generic equivalent. In conformance with State Education Law, which intends that patients receive the lower cost alternative, brand name drugs included in this program:

- Do not require "Dispense as Written" (DAW) or "Brand Medically Necessary" on the prescription;
- Have a generic co-payment;
- Are paid at the Brand Name Drug reimbursement rate or usual and customary price, whichever is lower (SMAC/FUL are not applied);
- Do not require a new prescription if the drug is removed from this program.

Once it is determined that the generic drug is more cost-effective than the brand name equivalent, the prior authorization requirement will be removed for the generic drug. In SFY 15/16, the savings achieved by this program was \$22,979,950.

Brand name drugs that were subject to this program at the end of this fiscal year include:

Abilify tablet	Combivir	Mepron	Tobradex suspension
Adderall XR	Copaxone 20 mg/mL SubQ	Myfortic	Tricor
Aggrenox	Diastat	Niaspan	Trilipix
Aldara	Epivir HBV tablet	Patanase	Trizivir
Alphagan P 0.15%	Exelon Patch	Protopic	Valcyte
Astepro	Focalin XR 5mg, 10 mg, 15mg, 20mg, 30mg, 40mg	Pulmicort Respules	Wellbutrin
Baraclude	Gabitril 2mg, 4mg	Soriatane	Xeloda
Catapres-TTS	Gleevec	Tegretol suspension	Xenazine
Cellcept suspension	Hepsera	Tegretol XR	

# The Preferred Diabetic Supply Program (PDSP) Diabetic Supply Program

As a result of legislation passed in 2008, the New York State Medicaid Program implemented, on October 1, 2009, the Preferred Diabetic Supply Program (PDSP). The PDSP was originally established for fee-for-service, Medicaid Managed Care and Family Health Plus members. The program does not include Medicare/Medicaid dually enrolled members. The PDSP covers a wide variety of blood glucose monitors and test strips provided by pharmacies and durable medical equipment providers through use of a preferred supply list (PSL). In SFY 15/16, a total of 68,915 diabetic supply claims were processed achieving gross savings through manufacturer rebates of \$5,942,003. In the prior SFY, 89,400 diabetic supply claims were processed with a gross savings of \$7,516,554. Diabetic supply rebates by county have been included in Appendix 10.

### The Prior Authorization Process

**Prior Authorization (PA)** is a management tool that seeks to assure that medically necessary cost-effective drug therapy is prescribed. All drugs with prior authorization requirements continue to be available to Medicaid members. Prior authorizations may occur automatically, through a comparison of claims to pre-determined criteria at the point-of-service (POS), or they may be requested by the prescriber's office by phone or fax or can be requested through PAXpress®, a Web based tool. The automated PA system utilizes pharmacy and medical claims

data to process a request against pre-defined criteria to determine if the patient meets clinical criteria requirements instantaneously. The ability to incorporate pharmacy and medical claims data into criteria allows for the creation of more clinically driven criteria to help ensure appropriate medication utilization, and does so without prescriber involvement. Since the implementation of the automated prior authorization system on December 29, 2011, approximately 5.09 million electronic prior authorizations have been issued without prescriber involvement. Over 91 percent of all prior authorizations issued this fiscal year were issued electronically. For SFY 15/16, 1,328,847 automated PAs were issued without prescriber involvement. The reduction in the need for prescriber involvement results in prescribers being able to devote more time to patient care that would have otherwise been spent on the phone or completing paperwork.

The Clinical Call Center (CCC), operated by Magellan Medicaid Administration is available twenty-four (24) hours a day, seven (7) days a week. Performance is monitored closely by the DOH to ensure appropriate and timely response to prescriber and pharmacy requests, and to ensure that members are afforded the protections required by law.

For this fiscal year, the CCC received approximately 197,645 phone requests and 109,226 fax requests for prior authorization under the PDP and CDRP. Nearly all phone requests (99.96 percent) were completed during the initial call. In addition, the CCC provided approximately 85,839 callers with general information or technical assistance with the PA process and identified and referred two potential instances of fraud and/or abuse to the Department. The CCC and quality assurance team continued to aid the DOH, Office of Medicaid Inspector General (OMIG) and Office of the Attorney General (OAG) in collecting data related to suspected fraud cases.

Medicaid enrolled prescribers can also initiate prior authorization requests using a web-based application. PAXpress<sup>®</sup> is a web-based pharmacy PA request/response application that is accessible through eMedNY.

### Preferred Drug Program (PDP) Prior Authorization Process

Under the PDP, prescribers or their authorized agents (such as a nurse or office staff), contact the CCC by phone or fax to present medical justification for non-preferred drugs. The criteria used by the CCC staff to evaluate a request for a non-preferred drug is set forth in legislation and consists of the following:

- The preferred drug has been tried by the patient and has failed to produce the desired health outcomes;
- The patient has tried the preferred drug and has experienced unacceptable side effects:
- The patient has been stabilized on a non-preferred drug and transition to the preferred drug would be medically contraindicated.
- Other clinical indications for the use of the non-preferred drug, which shall include consideration of the medical needs of special populations, including children, the elderly, the chronically ill, persons with mental health conditions, and persons affected by HIV/AIDS.

In general, prescribers initially speak with a Certified Pharmacy Technician (CPT) when requesting authorization for a non-preferred drug or a drug requiring prior authorization due to FQD, diagnosis or step therapy requirements. If the responses to the clinical criteria support the PA request, a PA is issued by the CPT. In the event the request does not meet the criteria; the call is referred to a pharmacist so that the prescriber may provide additional information that would support the use of the non-preferred drug. If, after that discussion, the clinical criteria are met, a PA is issued. However, as required by legislation, when a prescriber maintains that the use of the non-preferred drug is necessary, despite not meeting the clinical criteria, the

prescriber's determination prevails, and PA is granted. This occurred in 22.02 percent of the PDP PAs processed in SFY 15/16. Examples of PA requests where providers have utilized the prescriber prevails clause includes PA requests for:

- Second generation antipsychotics: patient does not meet diagnosis/age requirements in clinical criteria:
- Hepatitis C agents: prescriber does not provide clinical justification that would support the use of the preferred agent; and
- Inhaled antibiotics: prescriber is not familiar with the preferred agents and does not wish to try them.

### Clinical Drug Review Program (CDRP) Prior Authorization Process

Initially, the prescriber speaks with a CPT when requesting authorization. For select CDRP medications, only the prescriber who orders a CDRP drug can initiate the PA process. If, during the discussion, the clinical criteria for approval are not met, the request is referred to a pharmacist so that the prescriber may provide additional information to support the use of the drug. At the prescriber's request, a physician peer review may take place. In SFY 15/16, there were 56 physician peer reviews completed, however, consistent with last year, there were no denials rendered. Unlike the PDP which allows the prescriber to prevail, the CDRP legislation allows for a denial where there is substantial evidence of fraud or abuse. Under current statute, requests may not be denied for lack of medical necessity.

### III. Outreach and Education

Outreach and education efforts focus on ensuring that providers and members are informed about Medicaid's pharmacy PA programs and are kept up to date on program changes.

During the fiscal year, changes to the PDP occurred through the re-review of existing classes and addition of new drug classes. With each change, prescribers and pharmacies were notified in advance when the Preferred Drug List (PDL) was changing and the PA requirements that would apply to newly non-preferred and CDRP drugs. Notification was achieved via electronic notification and the Medicaid Update (a monthly Medicaid provider communication). The PDP website (newyork.fhsc.com) is another venue for information, offering easy access for prescribers, pharmacists, members and other interested parties (Appendix 7). Brochures for members are available on-line and in a number of languages including Bosnian, Chinese, Spanish, Yiddish and Haitian Creole (Appendix 6).

# IV. Prescriber, Pharmacy, and Patient Satisfaction

### **Complaints**

Complaints may be received through a variety of sources including by mail or email, through the Clinical Call Center (CCC) or Medicaid Helpline. When such calls are received they are referred to the DOH Medicaid pharmacy staff where direct assistance is provided. Seven complaints about the PDP and CDRP were received during this fiscal year, primarily via phone calls. Ten fewer complaints were received this SFY than were received the previous year. All complaints received (particularly those that are logged as "Other") are shared with the Quality Assurance Group (QAG) for review/follow-up and are used as a means for quality analysis/trending of data. Data are used as part of a continuous quality improvement process to ensure appropriate and timely response to complaints and to address opportunities for improvement. These complaints are listed below by the category in which they were logged.

Customer Service Pharmacy	3
Other	2
PA/Utilization Management Issue	1
PDL Criteria	1
Total	7

The DOH Medicaid pharmacy staff individually addresses issues related to policy. These inquiries are also used to identify providers who may need additional program education.

Patient reaction to the PDP remains positive. Medicaid's Helpline for members received 31 calls on this topic, but when such calls are received, they are referred to the DOH Medicaid pharmacy staff, which provides direct assistance to the member and/or their providers.

# V. Outcomes and Cost Savings

### Preferred Drug Program

Under the Medicaid Drug Rebate Program created by the Omnibus Reconciliation Act of 1990 (OBRA), drug manufacturers are required to enter into rebate agreements with the Centers for Medicare and Medicaid Services (CMS), for drug products reimbursed by Medicaid. Medicaid programs must cover all outpatient drugs of a manufacturer that signs a national rebate agreement. Many Medicaid programs, including New York's, use a PDP to collect supplemental rebates from manufacturers when their drugs are designated as preferred within the drug class.

To receive supplemental rebates, New York State joined the National Medicaid Pooling Initiative (NMPI). Additionally, the New York State Direct Contracting Program (SDC) enables access to rebates for manufactures that do not participate in NMPI. Both programs are administered by Magellan Medicaid Administration. New York is among 11 states that currently participate in the NMPI. Others include Alaska, Kentucky, Michigan, Minnesota, Montana, New Hampshire, Rhode Island, South Carolina, North Carolina and the District of Columbia. At the end of the fiscal year the NMPI includes more than 90 participating manufacturers and has approximately 5.5 million member lives.

Contracts with manufacturers have a three-year net price guarantee; net prices may decrease during the period, but they may not increase. Rebate amounts are based on the Wholesale Acquisition Cost (WAC) for each individual drug. Each Participating State in the NMPI program maintains its own P&TC or DURB and the ability to designate a drug as preferred or non-preferred.

The Medicaid Fee-for-Service program paid approximately 11.9 million paid pharmacy claims in SFY 15/16. Of these, 36 percent were for a drug that fell within one of the classes of drugs on the PDP. Of the drugs subject to the PDP, at the end of the fiscal year 66.4 percent of claims were for drugs that did not require prior authorization. The remaining 33.6 percent of claims were for drugs that required prior authorization. These percentages are attributable to the wide selection of preferred drugs within a class, prescriber familiarity with the Medicaid PDP and education efforts. Success is further supported by the pharmacy provider community in advising prescribers of preferred drug choices. There were 157,259 prior authorizations administered for <u>all</u> pharmacy programs.

Under the PDP, the highest volume of requests for prior authorizations during SFY 15/16 were for the following drug classes: short-acting opioids (20 percent), used to treat moderate to severe pain; second generation antipsychotics (19 percent), primarily used to treat mental health illnesses such as schizophrenia and bipolar disorder; CNS Stimulants (7 percent), primarily used to treat ADHD; Proton Pump Inhibitors (6 percent), used to treat acid reflux; and second generation anticonvulsants (5 percent), used primarily to treat seizure disorders. Requests for prior authorization for Hepatitis C Agents made up 1.3 percent of prior authorizations for this fiscal year.

Consistent with the experience in SFY 14/15, primary indicators for PDP PA requests to prescribe a non-preferred drug include treatment failure on preferred medication, contraindications preventing transition to preferred medications and adverse reactions to preferred medications. Overall, after consultation with CCC staff, 2.6 percent of the total requests resulted in the prescriber agreeing to use the preferred drug in lieu of a non-preferred drug. The CCC representatives have continued to promote the use of preferred agents as clinically appropriate, attributing to the relative changes observed. Total PDP savings are

calculated by combining the sum of supplemental rebates invoiced with the savings associated with market share shift to less expensive products within each therapeutic drug class. For SFY 15/16, total PDP savings were approximately \$12.8 million. As in the previous SFY, the FDA approval and FFS Medicaid coverage of new Hepatitis C Direct Acting Antivirals shortly before and during this time period significantly increased cost and negatively impacted total PDP savings. When these costly new drugs entered the market, they gained market share at the expense of older, lower cost products in their respective class. As a result, market shift savings over the period was negative. Appendix 10 lists the program's cost avoidance by county.

Market Shift Cost Avoidance is the difference between what was actually paid and what would have been paid without a Prescription Drug Plan (PDP). This will be negative if the net cost of the preferred agents (not including supplemental rebates) is higher than the net cost of the non-preferred agents. When costly new drugs enter the market, they may pick up market share at the expense of lower cost products. To that point, a negative market shift is not necessarily reflective of a poor PDL performance, because without the PDL the negative shift in market share towards the high-cost products would have been higher.

### Outcomes and Cost Savings – Clinical Drug Review Program (CDRP)

In this fiscal year, a total of 9,324 requests were approved for PA of drugs under the CDRP as follows:

Anabolic Steroids: 596

• CNS Stimulants (18 or Older): 5680

• Fentanyl Mucosal Agents: 52

• Growth Hormones (21 or Older): 14

• Immunomodulators: Topical: 282

• Lidoderm<sup>®</sup>: 636

• Phosphodiesterase type-5 (PDE-5) Inhibitors for PAH: 118

Regranex<sup>®</sup>: 14

Serostim<sup>®</sup>: 4

• Synagis<sup>®</sup>: 526

• Truvada<sup>®</sup>: 1155

• **Xyrem**<sup>®</sup>: 12

Oxazolidinone Antibiotics®: 235

All CDRP requests were authorized using the criteria in current statute, which allows a denial only based on substantial evidence of fraud and abuse. It is difficult to obtain evidence or documentation during a phone call, that would serve to support such a denial. However, if statute allowed denial based on medical necessity, 4 percent of requests would have been denied. This suggests that although the program has a strong sentinel effect, helping to ensure appropriate prescribing practices and protect patient safety, opportunities exist to enhance the program further.

### VI. Conclusion

The tenth full fiscal year of operation of the PDP, and CDRP, proceeded smoothly. Results continue to show that the PDP and CDRP programs are effective in assuring access to high quality, cost effective medications and have resulted in significant program savings, while promoting access to medically necessary drugs for Medicaid members.

In SFY 15/16, the DURB re-reviewed 28 classes of drugs in the PDP to include drugs recently approved by the FDA and newly available clinical and financial information. Five new drug classes were reviewed for inclusion on the PDP. By the end of the SFY there were a total of 108 drug classes subject to the PDP. No new drugs were recommended for inclusion into the CDRP by the DUR Board in this fiscal year.

Technological advancements including audiocasts of DURB meetings and email notification to interested parties regarding PDL changes, have ensured the transparency of the PDP and CDRP process.

Providers continue to receive notification of PDL revisions through email distribution lists, website postings and Medicaid Update article publications.

Since October 2011, members in mainstream Medicaid managed care plans receive their pharmacy benefit through their plans. This change explains the variance in rebates from this year compared to years prior to October 2011. The Medicaid FFS PDP continues to provide value to members that remain in FFS through the use of a preferred drug list which promotes clinically appropriate drug utilization, while also reducing costs.

The Pharmacy Prior Authorization programs continue to be monitored closely by DOH staff. An annual review of the NMPI and SDC supplemental invoice process by an independent consultant, is conducted to ensure appropriate protocol and accounting is maintained. Complaints are tracked to ensure appropriate resolution, and feedback from complaints is evaluated for potential enhancements to the process.

# VII. Appendices

# 1 – Legislation: Part C, Section 10 of Chapter 58 of the Laws of 2005

#### ARTICLE 2-A

#### PRESCRIPTION DRUGS

- Section 270. Definitions.
  - 272. Preferred drug program.
  - 273. Preferred drug program prior authorization.
  - 274. Clinical drug review program.
  - 275. Applicability of prior authorization to EPIC.
  - 276. Education and outreach.
  - 277. Review and reports.
  - 280. Prescription drug discount program.
- § 270. Definitions. As used in this article, unless the context clearly requires otherwise:
  - 1. "Administrator" means an entity with which the commissioner contracts for the purpose of administering elements of the preferred drug program, as established under section two hundred seventy-two of this article or the clinical drug review program established under section two hundred seventy-four of this article.
    - 2. "Board" shall mean the drug utilization review board.
  - 3. "Clinical drug review program" means the clinical drug review program created by section two hundred seventy-four of this article.
  - 4. "Emergency condition" means a medical or behavioral condition as determined by the prescriber or pharmacists, the onset of which is sudden, that manifests itself by symptoms of sufficient severity, including severe pain, and for which delay in beginning treatment prescribed by the patient's health care practitioner would result in:
  - (a) placing the health or safety of the person afflicted with such condition or other person or persons in serious jeopardy;
    - (b) serious impairment to such person's bodily functions;
    - (c) serious dysfunction of any bodily organ or part of such person;
    - (d) serious disfigurement of such person; or
    - (e) severe discomfort.
  - 5. "Non preferred drug" means a prescription drug that is included in the preferred drug program and is not one of the drugs on the preferred drug list because it is either: (a) in a therapeutic class that is included in the preferred drug program and is not one of the drugs on the preferred drug list in that class or (b) manufactured by a pharmaceutical manufacturer with whom the commissioner is negotiating or has negotiated a manufacturer agreement and is not a preferred drug under a manufacturer agreement.
  - 6. "Panel" means the elderly pharmaceutical insurance coverage panel established pursuant to section two hundred forty-four of the elder law.
  - 7. "Preferred drug" means a prescription drug that is either (a) in a therapeutic class that is included in the preferred drug program and is one of the drugs on the preferred drug list in that class or (b) a preferred drug under a manufacturer agreement.
  - 8. "Preferred drug program" means the preferred drug program established under section two hundred seventy-two of this article.
  - 9. "Prescription drug" or "drug" means a drug defined in subdivision seven of section sixty-eight hundred two of the education law, for which a prescription is required under the federal food, drug and cosmetic act. Any drug that does not require a prescription under such act, but which would otherwise meet the criteria under this article for inclusion on the preferred drug list may be added to the preferred drug list under this article; and, if so included, shall be considered to be a

prescription drug for purposes of this article; provided that it shall be eligible for reimbursement under a state public health plan when ordered by a prescriber authorized to prescribe under the state public health plan and the prescription is subject to the applicable provisions of this article and paragraph (a) of subdivision four of section three hundred sixty-five-a of the social services law.

- 10. "Prior authorization" means a process requiring the prescriber or the dispenser to verify with the applicable state public health plan or its authorized agent that the drug is appropriate for the needs of the specific patient.
- 11. "State public health plan" means the medical assistance program established by title eleven of article five of the social services law (referred to in this article as "Medicaid"), the elderly pharmaceutical insurance coverage program established by title three of article two of the elder law (referred to in this article as "EPIC"), and the family health plus program established by section three hundred sixty-nine-ee of the social services law to the extent that section provides that the program shall be subject to this article.
- 12. "Supplemental rebate" means a supplemental rebate under subdivision eleven of section two hundred seventy-two of this article.
- 13. "Therapeutic class" means a group of prescription drugs that produce a particular intended clinical outcome and are grouped together as a therapeutic class by the pharmacy and therapeutics committee.
- 14. "Manufacturer agreement" means an agreement between the commissioner and a pharmaceutical manufacturer under paragraph (b) of subdivision eleven of section two hundred seventy-two of this article.
- § 272. Preferred drug program. 1. There is hereby established a preferred drug program to promote access to the most effective prescription drugs while reducing the cost of prescription drugs for persons in state public health plans.
  - 2. When a prescriber prescribes a non-preferred drug, state public health plan reimbursement shall be denied unless prior authorization is obtained, unless no prior authorization is required under this article.
  - 3. The commissioner shall establish performance standards for the program that, at a minimum, ensure that the preferred drug program and the clinical drug review program provide sufficient technical support and timely responses to consumers, prescribers and pharmacists.
  - 4. Notwithstanding any other provision of law to the contrary, no preferred drug program or prior authorization requirement for prescription drugs, except as created by this article, paragraph (a-1) or (a-2) of subdivision four of section three hundred sixty-five-a of the social services law, paragraph (g) of subdivision two of section three hundred sixty-five-a of the social services law, subdivision one of section two hundred forty-one of the elder law and shall apply to the state public health plans.
  - 5. The drug utilization review board shall consider and make recommendations to the commissioner for the adoption of a preferred drug program. (a) In developing the preferred drug program, the board shall, without limitation: (i) identify therapeutic classes or drugs to be included in the preferred drug program; (ii) identify preferred drugs in each of the chosen therapeutic classes; (iii) evaluate the clinical effectiveness and safety of drugs considering the latest peer-reviewed research and may consider studies submitted to the federal food and drug administration in connection with its drug approval system; (iv) consider the potential impact on patient care and the potential fiscal impact that may result from making such a therapeutic class subject to prior authorization; and (v) consider the potential impact of the preferred drug program on the health of special populations such as

children, the elderly, the chronically ill, persons with  ${\tt HIV/AIDS}$  and persons with mental health conditions.

- (b) In developing the preferred drug program, the board may consider preferred drug programs or evidence based research operated or conducted by or for other state governments, the federal government, or multi-state coalitions. Notwithstanding any inconsistent provision of section one hundred twelve or article eleven of the state finance law or section one hundred forty-two of the economic development law or any other law, the department may enter into contractual agreements with the Oregon Health and Science University Drug Effectiveness Review Project to provide technical and clinical support to the board and the department in researching and recommending drugs to be placed on the preferred drug list.
- (c) The board shall from time to time review all therapeutic classes included in the preferred drug program, and may recommend that the commissioner add or delete drugs or classes of drugs to or from the preferred drug program, subject to this subdivision.
- (d) The board shall establish procedures to promptly review prescription drugs newly approved by the federal food and drug administration.
- 6. The board shall recommend a procedure and criteria for the approval of non-preferred drugs as part of the prior authorization process. In developing these criteria, the board shall include consideration of the following:
- (a) the preferred drug has been tried by the patient and has failed to produce the desired health outcomes;
- (b) the patient has tried the preferred drug and has experienced unacceptable side effects;
- (c) the patient has been stabilized on a non-preferred drug and transition to the preferred drug would be medically contraindicated; and
- (d) other clinical indications for the use of the non-preferred drug, which shall include consideration of the medical needs of special populations, including children, the elderly, the chronically ill, persons with mental health conditions, and persons affected by HIV/AIDS.
- 7. The commissioner shall provide thirty days public notice on the department's website prior to any meeting of the board to develop recommendations concerning the preferred drug program. Such notice regarding meetings of the board shall include a description of the proposed therapeutic class to be reviewed, a listing of drug products in the therapeutic class, and the proposals to be considered by the board. The board shall allow interested parties a reasonable opportunity to make an oral presentation to the board related to the prior authorization of the therapeutic class to be reviewed. The board shall consider any information provided by any interested party, including, but not limited to, prescribers, dispensers, patients, consumers and manufacturers of the drug in developing their recommendations.
- 8. The commissioner shall provide notice of any recommendations developed by the board regarding the preferred drug program, at least five days before any final determination by the commissioner, by making such information available on the department's website. Such public notice may include: a summary of the deliberations of the board; a summary of the positions of those making public comments at meetings of the board; the response of the board to those comments, if any; and the findings and recommendations of the board.
- 9. Within ten days of a final determination regarding the preferred drug program, the commissioner shall provide public notice on the department's website of such determinations, including: the nature of the determination; and analysis of the impact of the commissioner's

determination on state public health plan populations and providers; and the projected fiscal impact to the state public health plan programs of the commissioner's determination.

- 10. The commissioner shall adopt a preferred drug program and amendments after considering the recommendations from the board and any comments received from prescribers, dispensers, patients, consumers and manufacturers of the drug.
- (a) The preferred drug list in any therapeutic class included in the preferred drug program shall be developed based initially on an evaluation of the clinical effectiveness, safety and patient outcomes, followed by consideration of the cost-effectiveness of the drugs.
- (b) In each therapeutic class included in the preferred drug program, the board shall determine whether there is one drug which is significantly more clinically effective and safe, and that drug shall be included on the preferred drug list without consideration of cost. If, among two or more drugs in a therapeutic class, the difference in clinical effectiveness and safety is not clinically significant, then cost effectiveness (including price and supplemental rebates) may also be considered in determining which drug or drugs shall be included on the preferred drug list.
- (c) In addition to drugs selected under paragraph (b) of this subdivision, any prescription drug in the therapeutic class, whose cost to the state public health plans (including net price and supplemental rebates) is equal to or less than the cost of another drug in the therapeutic class that is on the preferred drug list under paragraph (b) of this subdivision, may be selected to be on the preferred drug list, based on clinical effectiveness, safety and cost-effectiveness.
- (d) Notwithstanding any provision of this section to the contrary, the commissioner may designate therapeutic classes of drugs, including classes with only one drug, as all preferred prior to any review that may be conducted by the board pursuant to this section.
- 11. (a) The commissioner shall provide an opportunity for pharmaceutical manufacturers to provide supplemental rebates to the state public health plans for drugs within a therapeutic class; such supplemental rebates shall be taken into consideration by the board and the commissioner in determining the cost-effectiveness of drugs within a therapeutic class under the state public health plans.
- (b) The commissioner may designate a pharmaceutical manufacturer as one with whom the commissioner is negotiating or has negotiated a manufacturer agreement, and all of the drugs it manufactures or markets shall be included in the preferred drug program. The commissioner may negotiate directly with a pharmaceutical manufacturer for rebates relating to any or all of the drugs it manufactures or markets. A manufacturer agreement shall designate any or all of the drugs manufactured or marketed by the pharmaceutical manufacturer as being preferred or non preferred drugs. When a pharmaceutical manufacturer has been designated by the commissioner under this paragraph but the commissioner has not reached a manufacturer agreement with pharmaceutical manufacturer, then the commissioner may designate some or all of the drugs manufactured or marketed by the pharmaceutical manufacturer as non preferred drugs. However, notwithstanding this paragraph, any drug that is selected to be on the preferred drug list under paragraph (b) of subdivision ten of this section on grounds that it is significantly more clinically effective and safer than other drugs in its therapeutic class shall be a preferred drug.
- (c) Supplemental rebates under this subdivision shall be in addition to those required by applicable federal law and subdivision seven of section three hundred sixty-seven-a of the social services law. In order

- to be considered in connection with the preferred drug program, such supplemental rebates shall apply to the drug products dispensed under the Medicaid program and the EPIC program. The commissioner is prohibited from approving alternative rebate demonstrations, value added programs or guaranteed savings from other program benefits as a substitution for supplemental rebates.
- 13. The commissioner may implement all or a portion of the preferred drug program through contracts with administrators with expertise in management of pharmacy services, subject to applicable laws.
- 14. For a period of eighteen months, commencing with the date of enactment of this article, and without regard to the preferred drug program or the clinical drug review program requirements of this article, the commissioner is authorized to implement, or continue, a prior authorization requirement for a drug which may not be dispensed without a prescription as required by section sixty-eight hundred ten of the education law, for which there is a non-prescription version within the same drug class, or for which there is a comparable non-prescription version of the same drug. Any such prior authorization requirement shall be implemented in a manner that is consistent with the process employed by the commissioner for such authorizations as of one day prior to the date of enactment of this article. At the conclusion of the eighteen month period, any such drug or drug class shall be subject to the preferred drug program requirements of this article; provided, however, that the commissioner is authorized to immediately subject any such drug to prior authorization without regard to the provisions of subdivisions five through eleven of this section.
- § 273. Preferred drug program prior authorization. 1. For the purposes of this article, a prescription drug shall be considered to be not on the preferred drug list if it is a non preferred drug.
  - 2. The preferred drug program shall make available a twenty-four hour per day, seven days per week telephone call center that includes a toll-free telephone line and dedicated facsimile line to respond to requests for prior authorization. The call center shall include qualified health care professionals who shall be available to consult with prescribers concerning prescription drugs that are not on the preferred drug list. A prescriber seeking prior authorization shall consult with the program call line to reasonably present his or her justification for the prescription and give the program's qualified health care professional a reasonable opportunity to respond.
  - 3. (a) When a patient's health care provider prescribes a prescription drug that is not on the preferred drug list, the prescriber shall consult with the program to confirm that in his or her reasonable professional judgment, the patient's clinical condition is consistent with the criteria for approval of the non-preferred drug. Such criteria shall include:
  - (i) the preferred drug has been tried by the patient and has failed to produce the desired health outcomes;
  - (ii) the patient has tried the preferred drug and has experienced unacceptable side effects;
  - (iii) the patient has been stabilized on a non-preferred drug and transition to the preferred drug would be medically contraindicated; or
  - (iv) other clinical indications identified by the committee for the patient's use of the non-preferred drug, which shall include consideration of the medical needs of special populations, including children, elderly, chronically ill, persons with mental health conditions, and persons affected by HIV/AIDS.
    - (b) In the event that the patient does not meet the criteria in

- paragraph (a) of this subdivision, the prescriber may provide additional information to the program to justify the use of a prescription drug that is not on the preferred drug list. The program shall provide a reasonable opportunity for a prescriber to reasonably present his or her justification of prior authorization. If, after consultation with the program, the prescriber, in his or her reasonable professional judgment, determines that the use of a prescription drug that is not on the preferred drug list is warranted, the prescriber's determination shall be final.
- (c) If a prescriber meets the requirements of paragraph (a) or (b) of this subdivision, the prescriber shall be granted prior authorization under this section.
- (d) In the instance where a prior authorization determination is not completed within twenty-four hours of the original request, solely as the result of a failure of the program (whether by action or inaction), prior authorization shall be immediately and automatically granted with no further action by the prescriber and the prescriber shall be notified of this determination. In the instance where a prior authorization determination is not completed within twenty-four hours of the original request for any other reason, a seventy-two hour supply of the medication shall be approved by the program and the prescriber shall be notified of this determination.
- 4. When, in the judgment of the prescriber or the pharmacist, an emergency condition exists, and the prescriber or pharmacist notifies the program that an emergency condition exists, a seventy-two hour emergency supply of the drug prescribed shall be immediately authorized by the program.
- 5. In the event that a patient presents a prescription to a pharmacist for a prescription drug that is not on the preferred drug list and for which the prescriber has not obtained a prior authorization, the pharmacist shall, within a prompt period based on professional judgment, notify the prescriber. The prescriber shall, within a prompt period based on professional judgment, either seek prior authorization or shall contact the pharmacist and amend or cancel the prescription. The pharmacist shall, within a prompt period based on professional judgment, notify the patient when prior authorization has been obtained or denied or when the prescription has been amended or cancelled.
- 6. Once prior authorization of a prescription for a drug that is not on the preferred drug list is obtained, prior authorization shall not be required for any refill of the prescription.
- 7. No prior authorization under the preferred drug program shall be required when a prescriber prescribes a drug on the preferred drug list; provided, however, that the commissioner may identify such a drug for which prior authorization is required pursuant to the provisions of the clinical drug review program established under section two hundred seventy-four of this article.
- 8. The department shall monitor the prior authorization process for prescribing patterns which are suspected of endangering the health and safety of the patient or which demonstrate a likelihood of fraud or abuse. The department shall take any and all actions otherwise permitted by law to investigate such prescribing patterns, to take remedial action and to enforce applicable federal and state laws.
- 9. No prior authorization under the preferred drug program shall be required for any prescription under EPIC until the panel has made prior authorization applicable to EPIC under section two hundred seventy-five of this article.
- 10. Prior authorization shall not be required for an initial or renewal prescription for buprenorphine or injectable naltrexone for detoxification or maintenance treatment of opioid addiction unless the

prescription is for a non-preferred or non-formulary form of such drug as otherwise required by section 1927(k)(6) of the Social Security Act.

- 274. Clinical drug review program. 1. In addition to the preferred drug program established by this article, the commissioner may establish a clinical drug review program. The commissioner may, from time to time, require prior authorization under such program for prescription drugs or patterns of utilization under state public health plans. When a prescriber prescribes a drug which requires prior authorization under this section, state public health plan reimbursement shall be denied unless such prior authorization is obtained.
  - 2. The clinical drug review program shall make available a twenty-four hour per day, seven days per week response system.
  - 3. In establishing a prior authorization requirement for a drug under the clinical drug review program, the commissioner shall consider the following:
  - (a) whether the drug requires monitoring of prescribing protocols to protect both the long-term efficacy of the drug and the public health;
  - (b) the potential for, or a history of, overuse, abuse, drug diversion or illegal utilization; and
  - (c) the potential for, or a history of, utilization inconsistent with approved indications. Where the commissioner finds that a drug meets at least one of these criteria, in determining whether to make the drug subject to prior authorization under the clinical drug review program, the commissioner shall consider whether similarly effective alternatives are available for the same disease state and the effect of that availability or lack of availability.
  - 4. The commissioner shall obtain an evaluation of the factors set forth in subdivision three of this section and a recommendation as to the establishment of a prior authorization requirement for a drug under the clinical drug review program from the drug utilization review board. For this purpose, the commissioner and the board, as applicable, shall comply with the following meeting and notice processes established by this article:
  - (a) the open meetings law and freedom of information law provisions of subdivision six of section two hundred seventy-one of this article; and
  - (b) the public notice and interested party provisions of subdivisions seven, eight and nine of section two hundred seventy-two of this article.
  - 5. The board shall recommend a procedure and criteria for the approval of drugs subject to prior authorization under the clinical drug review program. Such criteria shall include the specific approved clinical indications for use of the drug.
  - 6. The commissioner shall identify a drug for which prior authorization is required, as well as the procedures and criteria for approval of use of the drug, under the clinical drug review program after considering the recommendations from the board and any comments received from prescribers, dispensers, consumers and manufacturers of the drug. In no event shall the prior authorization criteria for approval pursuant to this subdivision result in denial of the prior authorization request based on the relative cost of the drug subject to prior authorization.
  - 7. In the event that the patient does not meet the criteria for approval established by the commissioner in subdivision six of this section, the clinical drug review program shall provide a reasonable opportunity for a prescriber to reasonably present his or her justification for prior authorization. If, after consultation with the program, the prescriber, in his or her reasonable professional judgment, determines that the use of the prescription drug is warranted, the

prescriber's determination shall be final and prior authorization shall be granted under this section; provided, however, that prior authorization may be denied in cases where the department has substantial evidence that the prescriber or patient is engaged in fraud or abuse relating to the drug.

- 8. In the event that a patient presents a prescription to a pharmacist for a prescription drug that requires prior authorization under this section and for which prior authorization has not been obtained, the pharmacist shall, within a prompt period based on professional judgment, notify the prescriber. The prescriber shall, within a prompt period based on professional judgment, either seek prior authorization or shall contact the pharmacist and amend or cancel the prescription. The pharmacist shall, within a prompt period based on professional judgment, notify the patient when prior authorization has been obtained or denied or when the prescription has been amended or cancelled.
- 9. In the instance where a prior authorization determination is not completed within twenty-four hours of the original request solely as the result of a failure of the program (whether by action or inaction), prior authorization shall be immediately and automatically granted without further action by the prescriber and the prescriber shall be notified of this determination. In the instance where a prior authorization determination is not completed within twenty-four hours of the original request for any other reason, a seventy-two hour supply of the medication will be approved by the program and the prescriber shall be notified of the determination.
- 10. When, in the judgment of the prescriber or the pharmacist, an emergency condition exists, and the prescriber or pharmacist notifies the program to confirm that such an emergency condition exists, a seventy-two hour emergency supply of the drug prescribed shall be immediately authorized by the program.
- 11. The department or the panel shall monitor the prior authorization process for prescribing patterns which are suspected of endangering the health and safety of the patient or which demonstrate a likelihood of fraud or abuse. The department or the panel shall take any and all actions otherwise permitted by law to investigate such prescribing patterns, to take remedial action and to enforce applicable federal and state laws.
- 12. The commissioner may implement all or a portion of the clinical drug review program through contracts with administrators with expertise in management of pharmacy services, subject to applicable laws.
- 13. No prior authorization under the clinical drug review program shall be required for any prescription under EPIC until the commissioner has made prior authorization applicable to EPIC under section two hundred seventy-five of this article.
- 14. For the period of eighteen months, commencing with the date of enactment of this article, the commissioner is authorized to continue prior authorization requirements for prescription drugs subject to prior authorization as of one day prior to the enactment of this article and which are not described in subdivision fourteen of section two hundred seventy-two of this article. At the conclusion of the eighteen month period, any such drug shall be subject to the clinical drug review program requirements of this section; provided, however, that the commissioner is authorized to immediately subject any such drug to prior authorization without regard to the provisions of subdivisions three through six of this section.
- § 275. Applicability of prior authorization to EPIC. The panel shall, no later than April first, two thousand eight, proceed to make prior authorization under the preferred drug program and the clinical review

drug program, under this article, applicable to prescriptions under EPIC. The panel shall take necessary actions consistent with this article to apply prior authorization under this article to EPIC. Upon determining that the necessary steps have been taken to apply prior authorization under this article to EPIC, the panel shall, with reasonable prior public notice, make prescriptions under EPIC subject to prior authorization under this article as of a specified date. If necessary, the panel may provide that such applicability take effect on separate dates for the preferred drug program and the clinical drug review program.

- § 276. Education and outreach. The department or the panel may conduct education and outreach programs for consumers and health care providers relating to the safe, therapeutic and cost-effective use of prescription drugs and appropriate treatment practices for containing prescription drug costs. The department or the panel shall provide information as to how prescribers, pharmacists, patients and other interested parties can obtain information regarding drugs included on the preferred drug list, whether any change has been made to the preferred drug list since it was last issued, and the process by which prior authorization may be obtained.
- § 277. Review and reports. 1. The commissioner, in consultation with the drug utilization review board, shall undertake periodic reviews, at least annually, of the preferred drug program which shall include consideration of:
  - (a) the volume of prior authorizations being handled, including data on the number and characteristics of prior authorization requests for particular prescription drugs;
  - (b) the quality of the program's responsiveness, including the quality of the administrator's responsiveness;
    - (c) complaints received from patients and providers;
  - (d) the savings attributable to the state, and to each county and the city of New York, due to the provisions of this article;
  - (e) the aggregate amount of supplemental rebates received in the previous fiscal year and in the current fiscal year, to date; and such amounts are to be broken out by fiscal year and by month;
  - (f) the education and outreach program established by section two hundred seventy-six of this article.
  - 2. The commissioner and the board shall, beginning March thirty-first, two thousand six and annually thereafter, submit a report to the governor and the legislature concerning each of the items subject to periodic review under subdivision one of this section.
  - 3. The commissioner and the board shall, beginning with the commencement of the preferred drug program and monthly thereafter, submit a report to the governor and the legislature concerning the amount of supplemental rebates received.

# 2 – Drug Utilization Review Board Membership (as of March 2016)

### Drug Utilization Review Board Membership

### DOH Designee - Chairperson

1. Jason Helgerson

### <u>Physicians</u>

- 2. Renante Ignacio, MD
- 3. Paula Panzer, MD
- 4. Asa Radix, MD
- 5. James Saperstone, MD
- 6. Christopher J. Murphy, MD
- 7. Vacancy

### Pharma cists

- 8. Lisa Anzisi, PharmD
- 9. Leigh Briscoe-Dwyer, PharmD
- 10. James R. Hopsicker, RPh, MBA
- 11. Michelle Rainka, PharmD
- 12. Tara M. Thomas, RPh, MBA
- 13. Vacancy

#### DUR Experts

- 14. Donna Chiefari, PharmD
- 15. Jadwiga Najib, PharmD

### Nurse Practioner/Midwife

16. Nancy Balkon, PhD, NP

### Consumers/Consumer Representatives

- 17. Marla Eglowstein, MD
- 18. John Wikiera
- 19. Vacancy

# 3 – Drug Classes in the Preferred Drug Program (as of March 2016)

The following table lists drug classes that were reviewed at the DURB during this fiscal year. Also included is the review date, the date the <u>PDL</u> was publicly posted, and the date some drugs within the class required PA.

DURB Meeting	Drug Class	Posting Date	Date PA Required
April 22, 2015	ACE INHIBITORS	May 26, 2015	June 25, 2015
September 1, 2015	ACE INHIBITOR COMBINATIONS	September 1, 2015	September 1, 2015
September 17, 2015	ACTINIC KERATOSIS AGENTS	October 27, 2015	November 19, 2015
September 17, 2015	ANABOLIC STEROIDS: TOPICAL	October 27, 2015	November 19, 2015
April 22, 2015	ANTIBIOTIC-STEROID COMBINATION: OPHTHALMIC	May 26, 2015	June 25, 2015
April 22, 2015	ANTICHOLINERGICS: RESPIRATORY	May 26, 2015	June 25, 2015
April 22, 2015	ANTICOAGULANTS: ORAL	May 26, 2015	June 25, 2015
April 22, 2015	ANTICONVULSANTS SECOND GENERATION	May 26, 2015	June 25, 2015
April 22, 2015	ANTIHISTAMINES: OPHTH	May 26, 2015	June 25, 2015
April 22, 2015	ANTIVIRALS: TOPICAL	May 26, 2015	June 25, 2015
September 1, 2015	ARB COMBINATIONS	September 1, 2015	September 1, 2015
April 22, 2015	BETA AGONISTS: LONG-ACTING	May 26, 2015	June 25, 2015
April 22, 2015	CNS STIMULANTS	May 26, 2015	June 25, 2015
April 22, 2015	ERYTHROPOIESIS STIMULATING AGENTS	May 26, 2015	June 25, 2015
April 22, 2015	GLP-1 RECEPTOR AGONISTS	May 26, 2015	June 25, 2015
April 22, 2015	IMMUNOMODULATORS: SYSTEMIC	May 26, 2015	June 25, 2015
September 17, 2015	ANTIBIOTICS, INHALED	October 27, 2015	November 19, 2015
September 17, 2015	INSULINS: RAPID-ACTING	October 27, 2015	November 19, 2015
April 22, 2015 MULTIPLE SCLEROSIS AGENTS		May 26, 2015	June 25, 2015
April 22, 2015	NIACIN DERIVATIVES	May 26, 2015	June 25, 2015
April 22, 2015	NSAIDS: PRESCRIPTION	May 26, 2015	June 25, 2015
April 22, 2015	OPIATES: LONG ACTING	May 26, 2015	June 25, 2015
April 22, 2015	OPIOID DEPENDENCE AGENTS	May 26, 2015	June 25, 2015
April 22, 2015	PHOSPHATE REGULATORS	May 26, 2015	June 25, 2015
September 17, 2015	PLATELET INHIBITORS	October 27, 2015	November 19, 2015
April 22, 2015	PSORIASIS AGENTS: TOPICAL	May 26, 2015	June 25, 2015
April 22, 2015	SEDATIVE HYPNOTICS	May 26, 2015	June 25, 2015
April 22, 2015	STEROIDS: INHALED	May 26, 2015	June 25, 2015
April 22, 2015	STEROIDS: TOPICAL - HIGH POTENCY	May 26, 2015	June 25, 2015
April 22, 2015	TRIG. LOWERING AGENTS	May 26, 2015	June 25, 2015
April 22, 2015	URINARY TRACT ANTISPASMODICS	May 26, 2015	June 25, 2015
September 17, 2015	ANTIPSYCHOTICS: SECOND GENERATION	October 27, 2015	November 19, 2015

# 4 – Preferred and Non-Preferred Drug List (as of March 2016)

Revised: March 15, 2016

Last Major Update: October 27, 2015

Last Update: February 21, 2013

Last Update: February 18, 2016

Last Update: November 6, 2014

Last Update: April 25, 2013

### New York State Medicaid Fee-For-Service Pharmacy Programs

### OVERVIEW OF CONTENTS

### Preferred Drug Program (PDP) (Pages 2-39)

The PDP promotes the use of less expensive, equally effective drugs when medically appropriate through a Preferred Drug List (PDL). All drugs currently covered by Fee-For-Service (FFS) Medicaid remain available under the PDP and the determination of preferred and non-preferred drugs does not prohibit a prescriber from obtaining any of the medications covered under Medicaid.

- Non-preferred drugs in these classes require prior authorization (PA), unless indicated otherwise.
- Preferred drugs that require prior authorization are indicated by footnote.
- Specific Clinical, Frequency/Quantity/Duration, Step Therapy criteria is listed in column at the right.

### Clinical Drug Review Program (CDRP) (Page 40)

The CDRP is aimed at ensuring specific drugs are utilized in a medically appropriate manner. Under the CDRP, certain drugs require prior authorization because there may be specific safety issues, public health concerns, the potential for fraud and abuse, or the potential for significant overuse and misuse.

#### Drug Utilization Review (DUR) Program (Pages 41-46)

Last Update: January 21, 2016 The DUR helps to ensure that prescriptions for outpatient drugs are appropriate, medically necessary, and not likely to result in adverse medical consequences. This program uses professional medical protocols and computer technology and claims processing to assist in the management of data regarding the prescribing and dispensing of prescriptions. Frequency/Quantity/Duration (F/Q/D) Program and Step Therapy parameters are implemented to ensure clinically appropriate and cost effective use of these drugs and drug classes.

### Brand Less Than Generic (BLTG) Program (Page 47)

The Brand Less Than Generic Program is a cost containment initiative which promotes the use of certain multi-source brand name drugs when the cost of the brand name drug is less expensive than the generic equivalent. This program is in conformance with State Education Law, which intends that patients receive the lower cost alternative.

### Mandatory Generic Drug Program (Page 48)

State law excludes Medicaid coverage of brand name drugs that have a Federal Food and Drug Administration (FDA) approved A-rated generic equivalent, unless a prior authorization is obtained. Drugs subject to the Preferred Drug Program (PDP), Clinical Drug Review Program (CDRP), and/or the Brand Less Than Generic (BLTG) Program are not subject to the Mandatory Generic Program.

### Dose Optimization Program (Page 49-51)

Dose optimization can reduce prescription costs by reducing the number of pills a patient needs to take each day. The Department has identified drugs to be included in this program, the majority of which have FDA approval for once-a-day dosing, have multiple strengths available in correlating increments at similar costs and are currently being utilized above the recommended dosing frequency.

For more information on the NYS Medicaid Pharmacy Programs: http://www.health.ny.gov/health\_care/medicaid/program/pharmacy.htm To contact the NYS Medicaid Pharmacy Clinical Call Center please call 1-877-309-9493 To download a copy of the Prior Authorization fax form go to https://newyork.fhsc.com/providers/PA\_forms.asp

### Revised: March 15, 2016

# NYS Medicaid Fee-For-Service Preferred Drug List

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<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs		Non-Prefe	erred Drugs	Prior Authorization/Coverage Parameters	
			I. AN	ALGESICS	
		Non-Stero	idal Anti-Inflamma	tory Drugs (NSAIDS) – Prescription	
ibuprofen indomethacin m eloxicam tablet naproxen naproxen EC	naproxen sodium piroxicam sulindac Voltaren <sup>©</sup> Gel	Anaprox® DS Arthrotec® Cambia™ Celebrex® ©© celecoxib ©© Daypro® diclofenac / misoprostol diclofenac sodium diclofenac sodium diclofenac sodium diclofenac topical soln diffunisal Duexis® etodolac etodolac ER Feldene® fenoprofen Flector® patch ffurbiprofen Indocin® indomethacin SR ketoprofen	ketoprofen SA ketorolac meclofenamate mefenamic acid meloxicam susp. Mobic® nabumetone Naprelan® Naprosyn® Naprosyn® EC naproxen CR oxaprozin Pennsaid® Ponstel® Sprix® Tivorbex® tolmetin Vimovo® Vivlodex™ Voltaren® XR Zipsor® Zorvolex®	CUNICAL CRITERIA (CC)  Celebrex® (celecoxib) — one of the following criteria will not require PA  Over the age of 65 years  Concurrent use of an anticoagulant agent  History of GI Bleed/Ulcer or Peptic Ulcer Disease	
			Opioid	Antagonists	
naloxone (syringe, vi nattrexone Narcan <sup>e</sup> (nasal spray ReVia <sup>e</sup>		Evzio <sup>9</sup>			
			Opioid Depen	dence Agents <sup>CC, FIQID</sup>	
buprenorphine Suboxone <sup>6</sup> (film)		Bunavail <sup>®</sup> buprenorphine/ nalox Zubsolv <sup>®</sup>	one (tablet)	CLINICAL CRITERIA (CC)      Medical necessity rationale for opioid therapy is required for patients on established buprenorphine therapy      QUANTITY LIMIT:      Buprenorphine sublingual (SL): Six (6) tablets dispensed as a 2-day supply, not to exceed 24 mg per day      Buprenorphine/naloxone tablet and film (Bunavail™, Suboxone®, Zubsolv®): Three (3) sublingual tablets or films per day, maximum of 90 tablets or films dispensed as a 30-day supply, not to exceed 24 mg-6 mg of Suboxone, or it's equivalent per day	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 / 19/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
		Long-Acting <sup>©</sup>
mog, 100 mog) f <sup>60</sup> f <sup>60</sup> morphine sulfate ER (capsule) f <sup>60</sup> f <sup>60</sup> morphine sulfate SR (tablet) f <sup>60</sup> f <sup>60</sup>	Avinza France Butrans Conzip St. France Conzip St. France Em beda ER Exalgo France fentanyl patch (37.5 mog, 62.5 mog, 87.5 mog) France hydromorphone ER Hysingla ER France Kadian France MS Contin France Nucynta ER St. France Oxycodone ER France Oxycodone ER France Oxycontin France Oxycontin France Oxycontin France Oxycontin ER St. France Ultram ER St. France Zohydro ER France	CLINICAL CRITERIA (CC)   Limited to a total offour (4) opioid prescriptions every 30 days; Exemption for diagnosis of cancer or sickle cell disease.   Medical necessity rationale for opioid therapy is required for patients on established buprenorphine therapy.   PA required for initiation of long-acting opioid therapy in opioid-naïve patients.   Exemption for diagnosis of cancer or sickle cell disease.   PA required for any additional long-acting opioid prescription for patients currently on long-acting opioid therapy.   Exemption for diagnosis of cancer or sickle cell disease.   PA required for initiation of opioid therapy in patients currently on benzo diazepine therapy.   Exemption for diagnosis of cancer or sickle cell disease.   PA required for initiation of opioid therapy in patients currently on benzo diazepine therapy.   STEPTHERAPY (ST)   Nevertial ER (tapentadol ER): Trial with tapentadol IR before tapentadol ER for patients who are naïve to a long-acting opioid.   Tramadol ER (tamadol naïve patients): Attempt treatment with IR formulations before the following ER formulations: Conzip. Tramadol ER, Ultram ER.   REGUENCY/QUANTITY/DURATION (F/Q/D) - Exemption for diagnosis of cancer or sickle cell disease.   Embeda (morphine ER/naitrexone):   Maximum 2 (two) units per day.   Nucynta ER (tapentadol ER):   Maximum 2 (two) units per day.   Nucynta ER (tapentadol ER):   Maximum 2 (two) units per day.   Output ER (tydrocodone ER):   Maximum 3 (the blest dispensed as a 30 day supply.   Zohydro ER (hydrocodone ER):   Maximum 1 (one) unit per day; 30 units per 30 days.   Hysingla MER (hydrocodone ER):   Maximum 1 (one) unit per day; 30 units per 30 days.   Hysingla MER (hydrocodone ER):   Maximum 1 (one) units per day, 60 units per 30 days.   Norphine ER (MS Contin 16mg, 30mg, 60mg only):   Maximum 3 (three) units per day, 00 units per 30 days.   Morphine ER (MS Contin 15mg, 30mg, 60mg only):   Maximum 4 units per day, up to 3 times a day, maximum 120 units per 30 days.   Morphine ER (MS Contin 15mg, 30mg, 60

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	Short-Acting <sup>©</sup>	
butalbital / APAP / caffeine / codeine FM/0 codeine FM/0 codeine / APAP FM/0 hydrocodone / APAP FM/0 hydrocodone / ibuprofen FM/0 Lottab® (elixir) FM/0 morphine IR FM/0 oxycodone / APAP FM/0 Reprexain® FM/0 tramadol Verdrocet™ FM/0 Xylon™ FM/0 Signature FM/0 Si	butalbital compound / code ine FN/0 butorphanol nasal spray Demerol® dihydrocodeine / aspirin / caffeine FN/0 dihydrocodeine / AP AP / caffeine FN/0 Dilaudid® FN/0 Fioricet® / code ine FN/0 Fiorinal® / code ine FN/0 hydromorphone FN/0 lbudone® FN/0 levorphanol meperidine Nucynta® ST, FN/0 Opana® FN/0 Oxaydo™ FN/0 oxycodone / aspirin FN/0 oxycodone / ibuprofen FN/0 pentazocine / naloxone Percocet® 2.5/325mg FN/0 Primlev™ FN/0 Roxicet® (solution) FN/0 Synalgos® DC FN/0 tram adol / AP AP FN/0 Tylenol® / code ine #4 FN/0 Ultracet® FN/0 Ultram® Vicoprofen® FN/0 Xartemis® XR FN/0 Xodol® FN/0 Zamicet® FN/0	CLINICAL CRITERIA (CC)  Limited to a total of four (4) opioid prescriptions every 30 days; Exemption for diagnosis of cancer or sickle cell disease  For opioid: Naïve patients - limited to a 15 days supply for all initial opioid prescriptions, except for patients with diagnosis of sickle cell disease or cancer  Medical necessity rationale for opioid therapy is required for patients on established buprenorphine therapy  P A required for initiation of opioid therapy in patients currently on benzodiazepine therapy  STEP THERAPY (ST)  Nucynta® (tapentadol IR) — Trial with tramadol and one (1) preferred opioid before tapenta dol immediate-release (IR)  FREQUENCY/QUANTITY/DURATION (F/Q/D)  Quantity Limits:  Nucynta® (tapentadol IR):  Maximum 6 (six) units per day; 180 units per 30 days  Nucynta® (tapentadol IR):  Maximum 6 (six) units per day; 180 units per 30 days  Morphine and congeners immediate-release (IR) non-combination not to exceed 500mg/day  Morphine and congeners immediate-release (IR) non-combination products (codeine, hydrom orphone, morphine, oxycodone, oxynorphone):  Maximum 6 (six) units per day, 180 (one hundred eighty) units per 30 (thirty) days  Additional/alternate parameters: To be applied to patients without a documented cancer or sickle cell diagnosis  Morphine and congeners immediate-release (IR) combination products maximum recomm ended:  acetaminophen (4 grams)  aspirin (5 grams)  or the FDA approved maximum opioid dosage as listed in the PI, whichever is less  Duration Limits:  90 days for patients without a diagnosis of cancer or sickle-cell disease.

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Prefe	erred Drugs	Non-Pref	erred Drugs	Prior Authorization/Coverage Parameters			
II. ANTI-INFECTIVES							
	Antibiotics – Inhaled 🥰 FQ/D						
Bethkis <sup>61</sup>	Kitabis <sup>e</sup> Pak¹	TOBI <sup>®</sup> Podhaler™ <sup>2</sup>	tobramycin solution <sup>2</sup>	Prior Authorization required for non-preferred agents effective 11 /19/2015			
Cayston <sup>61</sup>		TOBI <sup>®</sup> solution <sup>2</sup>		CLINICAL CRITERIA (CC)			
				Confirm diagnosis for the FDA-approved indication of cystic fibrosis			
				FREQUENCY/QUANTITY/DURATION (F/Q/D)			
				> Aztreonam (Cayston)			
				3 (three) ampules (3mL) per day			
				84 ampules (84 mL) per 56 day regimen (28 days on, 28 days off)			
				➤ Tobramydin inhalation solution (Bethkis, TOBI, Kitabis)			
				<ul> <li>2 (two) ampules (8 m L Bethkis, 10 m L TOBI, Kitabis Pak) per day</li> </ul>			
				<ul> <li>56 ampules (224 mL Bethkis, 280 mL TOBI, Kitabis Pak) per 56 day regimen (28 days on- 28 days off)</li> </ul>			
				> Tobramycin capsules with inhalation powder (TOBI Podhaler)			
				8 capsules per day 224 capsules per 56 day regimen (28 days on-28 days off)			
			Anti-Fungals – O	ral for Onychomycosis			
griseofulvin (suspension) Grifulvin V <sup>6</sup> (tablet)							
griseofulvin ultramicronized		Gris-PEG <sup>●</sup>					
terbinafine (tablet)	)	griseoful vin micronizo	ed (tablet)				
		itraconazole					
		Lamisil <sup>e</sup> (tablet)					
		Om nel <sup>®</sup>					
		Sporanox <sup>6</sup>					
			Anti-V	firals – Oral			
acydovir		famciclovir	Valtrex <b>®</b>				
valacyclovir		Fam vir <sup>€</sup>	Zovirax <sup>®</sup>				
			Cephalosporin	s – Third Generation			
œfdinir	cefpodoxime	Cedax <sup>●</sup>	ceftibuten				
cefixime	Suprax <sup>6</sup>	ce fditoren					
			Fluoroqui	inolones – Oral			
Cipro <sup>6</sup> (suspensio	ın)	Avelox <sup>6</sup>	Le vaquin <sup>6</sup>				
ciprofloxacin (suspension, tablet)  Avelox ABC Pack levo floxacin (solution)		levo floxacin (solution)					
levofloxacin (table		Cipro <sup>e</sup> (tablet)	m oxifloxa cin				
	~	Cipro <sup>®</sup> XR	ofloxacin (tablet)				
		ciprofloxacin ER					

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Revised: March 15, 2016

Preferred Drugs		Non-Pre	ferred Drugs	Prior Authorization/Coverage Parameters
			Hepat	itis B Agents
Baraclude Hepsera adefovir dipivoxil lamiw Epivir-HBV Tyzeka entecavir		lamivudine 100m g		
		<u>'</u>	Hepatitis C Ag	ents – Injectable FRA'D
Pegasys <sup>®</sup>	PegIntron <sup>€</sup>	None		PRE QUENCY/QUANTITY/DURATION (F/Q/D)     PA required for the initial 14 weeks therapy to determine appropriate duration of therapy based on genotype, prior treatment and response, presence of cirrhosis, and HIV-coinfection.      Further documentation required for continuation of therapy at weeks 14 and 26.      After 12 weeks of therapy obtain a quantitative HCV RNA. Continuation is supported if undetectable HCV RNA or at least a 2 log decrease compared to baseline.      After 24 weeks of therapy obtain a HCV RNA. Continuation for genotype 1 and 4 is supported if undetectable HCV RNA.      Maximum duration of 48 weeks for:     ◆Treatment-naïve patients or prior relapsers with cirrhosis and HIV co-infection    ◆Prior non-responders (including prior partial and null responders) with or without cirrhosis and with or without HIV co-infection

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
		Hepatitis C Agents – Direct Acting Antivirals
ribavirin	Copegus <sup>6</sup>	CLINICAL CRITERIA (CC)
Viekira Pak™ <sup>cc</sup> , <sup>⊮</sup> Ω/0	Daklinza™ <sup>cc, F/Q/D</sup>	► Click here to access the Hepatitis C Worksheets with Clinical Criteria requirements  FREQUENCY/QUANTITY/DURATION (F/Q/D)
	Harvoni <sup>e cc, F/Q/D</sup>	> Daklinza™ (daolatasvir)
	l .	Quantity limit: Maxiumum 1 (one) unit per day; 28 units per 28 days
	Moderiba™	Duration limit: Initially 56 days, pending results of quantitative HCV RNA testing after 4 weeks of sofosbuvir/da clatas vir treatment
	Olysio <sup>ecc, Fa/0</sup> Rebetol <sup>e</sup>	<ul> <li>Maximum 12 consecutive weeks (all regimens given with sofosbuvir) over beneficiary lifetime, pending results of quantitative HCV RNA testing for patients with HCV genotypes 1 or 3 without cirrhosis; genotype 1 with compensated cirrhosis; genotype 3 with compensated</li> </ul>
		cirrhosis (with ribavirin); or genotypes 1 or 3 with decompensated cirrhosis or post-liver transplant (with ribavirin)  Harvoni™ (ledip asvir/sofosbuvir):
	Ribapak <sup>e</sup>	Quantity limit: Maxiumum 1 (one) unit per day; 28 units per 28 days
	Ribasphere **	Duration limit: Initially 56 days, pending results of quantitative HCV RNA testing after 4 weeks of ledipasvir/s of osbuvir treatment
	Sovaldi <sup>e co , F/Q/0</sup>	• Maximum 12 consecutive weeks over beneficiary lifetime, pending results of quantitative HCV RNA testing in patients with: genotype 1 and treatment-naïve without cirrhosis or with compensated cirrhosis; genotype 1 and treatment-experienced without cirrhosis; genotype 1
	Technivie <sup>TM CC, F/Q/D</sup>	with decompensated cirrhosis (given with ribavirin); genotypes 1 or 4 and post liver transplant without cirrhosis or with compensated
	Zepatier™ <sup>cc, ro/o</sup>	cirrhosis (given with ribavirin); or genotypes 4, 5, or 6 without cirrhosis or with compensated cirrhosis  Maximum 24 consecutive weeks over beneficiary lifetime, pending results of quantitative HCV RNA testing for: genotype 1 treatment-experienced patients with compensated cirrhosis
		> Ohysio™ (simeprevii):
		Quantity limit: Maximum 1 (one) unit per day, 28 units per 28 days
		<ul> <li>Duration limit: Initially 56 days, pending results of quantitative HCV RNA testing after 4 weeks of simeprevir treatment</li> <li>For Olysio plus peginterferon and rib avirin: maximum 12 consecutive weeks over beneficiarly lifetime for genotypes 1 and 4</li> </ul>
		◆ For Olysio plus Sovaldi; maximum 12 consecutive weeks for patients without cirrhosis and 24 weeks for patients with cirrhosis over
		ben eficiary lifetime for genotypes 1 and 4
		➤ Sovaldi™ (sofosbuvir):
		Quantity limit: Maximum 1 (one) unit per day, 28 units per 28 days
		Duration limit:  Maximum 12 consecutive weeks for: genotypes 1 or 4 if used with peginterferon and ribavirin; genotype 1 if used with Olysio in patients
		without cirrhosis; or genotype 2 if used with ribavirin
		<ul> <li>Maximum 24 consecutive weeks for: genotype 1 if used with Olysio in patients with cirrhosis; genotype 3 if used with ribavirin;</li> <li>Maximum up to 48 weeks in patients with hepatocellular carcinoma awaiting liver transplantation used in combination with ribavirin</li> </ul>
		➤ Technivie™ (ombitasvir/paritaprevii/ritonavir):
		Quantity limit: 56 units per 28 days; 2 units per dose pack; 1 dose pack per day
		Duration limit: Initially 56 days, pending results of quantitative HCV RNA testing after 4 weeks of Technivie treatment
		<ul> <li>Maximum 12 consecutive weeks over beneficiary lifetime, pending results of quantitative HCV RNA testing for patients with HCV</li> </ul>
		genotype 4 without cirrhosis  ➤ Viekira Pak™ (ombitasvir / paritaprevir / riton avir / das abuvir):
		Quantity limit: 112 units per 28 days; 4 units per blister pak; 1 blister pak per day
		<ul> <li>Duration limit: Initially 56 days, pending results of quantitative HCV RNA testing after 4 weeks of Viekira Pak treatment</li> <li>Maximum 12 consecutive weeks over beneficiary lifetime, pending results of quantitative HCV RNA testing for patients with HCV</li> </ul>
		genotype 1b and cirrhosis and for patients without cirrhosis [genotypes 1a or 1b, or unknown/mixed genotype 1 subtype(s)]  Maximum 24 consecutive weeks over beneficiary lifetime, pending results of quantitative HCV RNA testing for treatment experienced
		patients with HCV genoting a for unknown or mixed genoting a subtype(s)) who have cirrhosis
		> Zepatier™ (elbasvir/grazoprevir)
		Quantity limit: Maximum 1 (one) unit per day, 28 units per 28 days
		Duration limit: Initially 56 days, pending results of quantitative HCV RNA testing after 4 weeks of elbasvir/grazoprevir treatment
		♦ Maximum 12 consecutive weeks for members with the following: Genotype 1a and treatment-naïve or peginterferon + ribavirin-
		experienced and <u>without</u> baseline NSSA polymorphisms; genotype 1b treatment-naïve or peginterferon + ribavirin-experienced; or genotype 4 and treatment-naïve
		ribavirin-experienced and <u>with</u> baseline NSSA polymorphisms; or genotype 4 and peginterferon+ ribavirin-experienced
1 = Preferred as of 11/19/2	2015	8

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11/19/2015

Preferred Drugs	Non-Pre	ferred Drugs	Prior Authorization/Coverage Parameters
			etracyclines
dem edocydine doxycycline hydate minocydine (capsule) Morgidox <sup>®</sup> (capsule) tetracycline	Doryx <sup>® ST, F/Q /0</sup>		STEP THERAPY (ST)  Trial of a more cost effective doxycycline IR before progressing to doxycycline DR  FRE QUENCY/QUANTITY/DURATION (F/Q/D)  doxycycline DR (Doryx <sup>6</sup> )  Maximum 28 tablets/capsules per fill
		III. CA	RDIOVASCULAR
	А	ngiotensin Conver	ting Enzyme Inhibitors (ACEIs)
bena zepril lisino pril captopril rami pril enal april	Accupril <sup>®</sup> Altace <sup>®</sup> Epaned™ fosinopril Lotensin <sup>®</sup> Mavik <sup>®</sup> moexipril	perindopril Prinivil <sup>©</sup> quinapril trandolapril Univasc <sup>©</sup> Vasotec <sup>©</sup> Zestril <sup>©</sup>	
		ACE Inhi	bitor Combinations
bena zepril / amlodipine bena zepril / HCTZ captopril / HCTZ enalapril / HCTZ lisinopril / HCTZ Lotrel 6 moe xipril / HCTZ Tarka 6 trandolapril / verapamil ER	Accuretic <sup>©</sup> fosinopril/HCTZ Lotensin HCT <sup>©</sup> Prestalia <sup>©</sup>	quinapril/HCTZ Vaseretic <sup>©</sup> Zestoretic <sup>©</sup>	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Prefe	erred Drugs	Prior Authorization/Coverage Parameters
		Angiotensin Recep	otor Blockers (ARBs) <sup>ST</sup>
Diovan <sup>© 00</sup> valsartan Iosartan	Atacand <sup>©</sup> Avapro <sup>©</sup> Benicar <sup>© 10</sup> cande sartan Co zaar <sup>©</sup>	Edarbi™ eprosartan irbesartan Micardis <sup>© po</sup> telmisartan	DOSE OPTIMIZATION (DO)  See Dose Optimization Chart for affected drugs and strengths STEP THERAPY (ST)  Trial of a product containing ACE inhibitor prior to preferred ARB  Trial containing either an ACE inhibitor or ARB prior preferred direct renin inhibitor (DRI)
	00233		mbinations ST
Exforge HCT <sup>6</sup> losartan/ HCTZ valsartan/ amlodipine valsartan/ amlodipine / HCTZ valsartan/ HCTZ	Atacand HCT <sup>®</sup> Avalide <sup>®</sup> Azor <sup>®</sup> Benicar HCT <sup>® 00</sup> candesartan/ HCTZ Diovan HCT <sup>® 00</sup> E darbyclor <sup>™ 00</sup> Entrestor <sup>™</sup> Exforge <sup>® 00</sup> Hyzaar <sup>®</sup> irbesartan/ HCTZ Micardis HCT <sup>® 00</sup> telmisartan/ amlodipingtelmisartan/ HCTZ Tribenzor <sup>™</sup>	Đ	DOSE OPTIMIZATION (DO)  ➤ See Dose Optimization Chart for affected drugs and strengths  STEP THERAPY (ST)  ➤ Trial of product containing ACE Inhibitor prior to preferred ARB  ➤ Trial of product containing either ACE inhibitor or ARB prior to initiating DRI
		Beta	Blockers
atenolol carvedilol labetalol metoprolol succ. XL metoprolol tartrate propranolol (tablet)	acebutolol betaxolol bisoprolol Bystolic <sup>e po</sup> Coreg <sup>©</sup> Coreg CR <sup>© po</sup> Corgard <sup>©</sup> Inderal LA <sup>©</sup> InnoPran XL <sup>©</sup> Levatol <sup>©</sup>	Lopressor® nadolol pindolol propranolol (solution) propranolol ER/SA Sectral® Tenormin® timolol Toprol XL® DO Trandate® Zebeta®	DOSE OPTIMIZATION (DO) ➤ See Dose Optimization Chart for affected drugs and strengths

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs Non-F		Non-Prefe	rred Drugs	Prior Authorization/Coverage Parameters
			Beta Blo	ckers / Diuretics
atenolol/ chlorthalidone bisoprolol/ HCTZ propranolol/ HCTZ		Corzide <sup>®</sup> Dutoprol™ Lopressor HCT <sup>®</sup> m etoprolol tartrate/ HCTZ nadolo/ bendroflumethiazide Tenoretic <sup>®</sup> Ziac <sup>®</sup>		
			Calcium Channel E	Blockers (Dihydropyridine)
Afeditab CR e amlodipine felodipine ER isradipine	nicardipine HCI Nifedical XL <sup>®</sup> nifedipine nifedipine ER <i>I</i> SA	Adalat CC <sup>®</sup> nisoldipine Norvasc <sup>®</sup>	Procardia <sup>©</sup> Procardia XL <sup>©</sup> Sular <sup>©</sup>	
		•	Cholesterol A	Absorption Inhibitors
cholestyramine cholestyramine light Colestid <sup>®</sup> (tablet)	colestipol (tablet) Prevalite <sup>6</sup>	Colestid (granules) colestipol (granules) Questran <sup>6</sup>	Questran Light <sup>®</sup> Welchol <sup>®</sup> Zetia <sup>®</sup>	
			Direct Re	enin Inhibitors <sup>8T</sup>
Tekturna <sup>6</sup>	Tekturna HCT <sup>●</sup>	Amturnide <sup>™</sup>	Tekamlo <sup>™</sup>	STEP THERAPY (ST) Trial of product containing ACE Inhibitor prior to preferred ARB Trial of product containing either an ACE inhibitor or an ARB prior to initiating preferred DRI
	•	•	HMG-CoA Reduc	ctase Inhibitors (Statins)
atorvastatin lovastatin pravastatin	Simcor <sup>®</sup> sim vastatin	atorvastatin/amlodipine Caduet <sup>©</sup> Crestor <sup>© DO</sup> fluvastatin	Livalo <sup>©</sup> Pravachol <sup>©</sup> Vytorin <sup>©</sup>	DOSE OPTIMIZATION (DO) ➤ See Dose Optim ization Chart for affected drugs and strengths
	•	fluvastatin ER	Zocor <sup>6</sup>	» Padiustinas
Ni: <b>6</b> 00		niacin ER	Niaci	n Derivatives
Niaspan <sup>e po</sup>		macin E K		DOSE OPTIMIZATION (DO) ➤ See Dose Optim ization Chart for affected drugs and strengths

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11/19/2015

Pi	referred Drugs	Non-Prefe	erred Drugs	Prior Authorization/Coverage Parameters
	Pho sphodiest erase type-5			(PDE -5) Inhibitors for PAH CERP
Adcirca <sup>6</sup>	sildenafil	Revatio <sup>e</sup>		CLINICAL DRUG REVIEW PROGRAM (CDRP)
				➤ All prescriptions for <u>Addirca®</u> , <u>Revatio®and sildenafil</u> must have PA
				Prescribers are required to respond to a series of questions that identify prescriber, patient and reason for prescribing drug
				➤ Please be prepared to fax dinical documentation upon request
				➤ Prescriptions can be written for a 30-day supply with up to 5 refills
				➤ The <u>CDRP_Phosphodiesterase type-5 (PDE-5) Inhibitors for PAH_Prescriber Work sheet</u> provides step-by-step assistance in completing the prior authorization process
	Pulmonary Arterial Hyperte			nsion (PAH) Oral Agents, Other
Letairis <sup>6</sup>	Tradeer <sup>©</sup>	Adempas <sup>e</sup>	Orenitram <sup>6</sup>	FREQUENCY/QUANTITY/DURATION (F/Q/D)
		Opsumit <sup>e</sup>	Uptravi <sup>e rozo</sup>	> Uptravi <sup>®</sup> : 60 units per 30 days; 1 time titration packet
		•	Triglyceride	Lowering Agents
gem fibrozil		A⊓tara <sup>e</sup>		STEP THERAPY (ST)
Tricor <sup>e</sup>		fenofibrate		Lovaza® (omega-3-acid ethyl-esters) and Vascepa® (icosapent ethyl) — Trial of fibric acid
Trilipix <sup>©</sup>		fenofibric acid		derivative OR niadin prior to treatment with omega-3-adid ethyl-esters
		Fenoglide <sup>6</sup>		FREQUENCY/QUANTITY/DURATION (F/Q/D)
		Fibricor <sup>©</sup>		➤ <u>Lova za<sup>®</sup> (om eqa-3-acid ethyl-e sters) and Vascepa<sup>®</sup> (ico sapent ethyl)</u> – Required dosage
		Lipofen <sup>6</sup>		equal to 4 (four) units per day
		Lofibra <sup>®</sup>		
		Lopid <sup>®</sup>		
		Lovaza <sup>est, Horb</sup>	- F 10 10	
		omega-3 ethyl ester <sup>st</sup>	, Franc	
		Triglide <sup>6</sup>		
		Vascepa <sup>e st, F/Ω/0</sup>		

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
	IV. CENTRAL NERVOUS SYSTEM				
	Alzheimer's Agents				
donepezil 5mg , 10mg Exelon <sup>®</sup> (patch) galantamine galantamine ER memantine Namenda <sup>®</sup> rivastigmine (capsule)	Aricept <sup>®</sup> Namzaric <sup>® 22,87</sup> donepezil 23 mg rivastigmine (patch) Exelon <sup>®</sup> (capsule) Razadyne <sup>®</sup> Nam enda XR <sup>® 22,87</sup> Razadyne ER <sup>®</sup>	CLINICAL CRITERIA (CC)  ➤ Memantine extended-release containing products(Namenda XR™ and Namzaric™)— Require confirmation of diagnosis of dementia or Alzheimer's disease  STEP THERAPY (ST)  ➤ Memantine extended-release containing products (Namenda XR™ and Namzaric™)— Require trial with memantine immediate-release (Namenda <sup>®</sup> )			
	Anticonvulsants	- Second Generation			
felbamate gabapentin (capsule, solution) Gabitril (2mg, 4mg) lamotrigine (tablet) le vetiracetam le vetiracetam ER Lyrica (100 pm 1 m 200 topiramate (200 zonisamide	Banzel © 22 Felbatol © 25 Fycompa © 25 gabapentin (tablet) Gabitri © (12mg, 16mg) © 25 Keppra © 25 Keppra XR © 25 Lamictal © 25 Lamictal © 0DT Lamictal © 10T Lamictal © 10T Lamictal © 10T Lamictal © 10T Neurontin © 105 Info 02. ST Potiga © 125 Quidexy © XR Sabril © 125 Quidexy © XR Sabril © 125 Toparm ax © 125 Trokendi XR © 125 Zonegran © 125 Zonegran © 125	DOSE OPTIMIZATION (DO)			

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Pref	erred Drugs	Prior Authorization/Coverag	je Parameters
Antipsychotics – S		econd Generation <sup>CG</sup> <sup>ST</sup>		
Abilify <sup>© DO</sup> .1 clozapine Fanapt <sup>©</sup> Latuda <sup>© DO</sup> olanzapine (tablet) <sup>DO</sup> quetiapine <sup>FM270</sup> risperidone Saphris <sup>©</sup> Seroquel XR <sup>© DO</sup> .FM270 ziprasidone	aripiprazole aripiprazole ODT clozapine ODT Clozaril FazaClo Geodon Invega Invega ODT ODT DOT DOT DOT DOT DOT DOT DOT DOT	paliperidone ER FOOTD Rexulti FOOTD Resulti FOOTD Risperdal FOOTD Versadoz FOOTD Vraylar FOOTD Zyprexa FOOTD	DOSE OPTIMIZATION (DO)  See Dose Optimization Chart for affected drugs and CLINICAL CRITERIA (CC)  Clinical editing will allo wpatients currently stabilized receive that agent without PA  Require confirmation of FDA approved, compendials diagnosis is required for new prescriptions for all Seten Aripiprazole (Abilify). PA is not required when prefor schizophrenia as verified by Medicaid claims informed PA is required for initial prescription for beneficiaries minimum age as indicated below.  aripiprazole (Abilify)  asenapine (Saphris)  brexpiprazole (Rexutt)  cariprazine (Vraylar™)  clozapine (Clozaril, Fazaclo, Versacloz™)  iloperidone (Fanapt)  lurasidone HCI (Latuda)  olanzapine (Zyprexa)  paliperidone ER (Invega)  quetiapine fum. (Seroquel, Seroquel XR)  risperidone (Risperda)  ziprasidone HCI (Geodon)  Require confirmation of diagnosis that supports the cantipsychotic and a CNS Stimulant for patients < 18  STEP THERAPY (ST)  For all Second Generation Antipsychotics used the trin the absence of other psychiatric comorbidities, tria antidepressant agents is required  Trial of risperidone prior to paliperidone (Invega) the free QUENCY/QUANTITY/DURATION (F/Q/D)  cariprazine (Vraylar) 3mg, 4.5mg, 6mg: Maximum 1  cariprazine (Vraylar) 1.5mg: Maximum 2 (two) units paliperidone ER (Invega) 1.5mg, 3mg, 9mg tablets:  paliperidone ER (Invega) 6mg tablets: Maximum 2 (quetiapine/Quetiapine extended-release (Seroquel Standing) quetiapine (Seroquel): Maximum 3 (three) units per maximum 800mg/day  quetiapine (Seroquel): Maximum 3 (three) units per	on a non-preferred agent to continue to supported, or Medicaid covered cond Generation Antipsychotics cribed for treatment of bipolar disorder mation younger than the drug-specific  6 years 10 years 18 years 18 years 18 years 19 years 10 years 10 years 10 years 10 years 20 years 21 years 21 years 22 years 23 years 24 years 25 years 26 years 27 years 28 years 29 years 20 years 20 years 21 years 21 years 22 years 33 years 34 years 25 years 36 years 26 years 27 years 28 years 29 years 20 years 20 years 31 years 20 years 40 years 41 years 42 years 43 years 44 years 45 years 46 years 47 years 48 years 49 years 40 years 40 years 40 years 40 years 41 years 42 years 43 years 44 years 45 years 46 years 47 years 48 years 49 years 40 years 40 years 40 years 40 years 40 years 41 years 42 years 43 years 44 years 45 years 46 years 47 years 48 years 49 years 40
			<ul> <li>Seroquel XR* (150mg and 200mg): 1 (one) unit per</li> <li>Seroquel XR* (50mg, 300mg and 400mg): 2 (two) unit per</li> </ul>	day, 30 units per 30 days

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters	
Antipsychotics, Injectable			
Abilify Maintena®  fluphenazine decanoate  Haldol® decanoate  haloperidol decanoate  Invega Sustenna®  Invega Trinza™  Risperdal Consta®  Zyprexa Relprevv™			
	Benzodiaz	epines – Rectal	
Diastat <sup>6</sup> 2.5mg Diastat <sup>6</sup> AcuDial™	diazepam (rectal gel)		
	Carbamazepi	ine Derivatives <sup>©</sup>	
carbamazepine (chewable, tablet) carbamazepine ER (capsule) Epitol <sup>®</sup> Equetro <sup>®</sup> oxcarbazepine Tegretol <sup>®</sup> (suspension) Tegretol XR <sup>®</sup>	Aptiom <sup>6</sup> carbamazepine (suspension) carbamazepine XR (tablet) Carbatrol <sup>6</sup> Oxtellar XR <sup>6</sup> Tegretol <sup>6</sup> (tablet) Trileptal <sup>6</sup>	CLINICAL CRITERIA (CC) ➤ Clinical editing will allow patients currently stabilized on a non-preferred agent to continue to receive that agent without P A	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	Central Nervous System	(CNS) Stimulants C CORP FOOD
Adderall XR <sup>®</sup> Adderall XR <sup>®</sup> amphetamine salt combo immediate-release Daytrana <sup>®</sup> dexmethylphenidate dextroamphetamine (tablet) Focalin XR <sup>® 10</sup> Metadate <sup>®</sup> ER  Methylin <sup>®</sup> methylphenidate (tablet) methylphenidate ER (generic for Concerta <sup>®</sup> ) methylphenidate SR 10 mg, 20 mg (tablet) Quillivant XR <sup>®</sup> Vyvanse <sup>® 10</sup>	amphetamine salt combo extended-release Aptensio XRTM Concerta 00 Desoxyn Dexedrine dexmethylphenidate ER (generic Focalin XR 0) dexmethylphenidate ER (generic Focalin XR 0) dextroamphetamine ER dextroamphetamine (solution) Dyanavel XRTM Evekeo Focalin Metadate CD 00 methylphenidate CD (generic Metadate CD 0) methylphenidate ER (generic for Ritalin LA 0) methylphenidate (chewtablet, solution) modafinil Nuvigil 22, 00 Quillichew ER 1M Ritalin Ritalin LA 00 Zenzedi 0	CLINICAL CRITERIA (CC)  ➤ Confirm diagnosis for an FDA-approved or Compendia supported indication for beneficiaries less than 18 years of age.  ■ Prior authorization is required for initial prescriptions for stimulant therapy for beneficiaries less than 3 years of age  ■ Require confirm ation of diagnoses that support concurrent use of CNS Stimululant and Second Generation Antipsychotic agent  ➤ Patient-specific considerations for drug selection include treatment of excessive sleepiness associated with shift work sleep disorder or as an adjunct to standard treatment for obstructive sleep apnea.  CLINICAL DRUG RE VIEW PROGRAM (CDRP)  ➤ For patients 18 years of age and older:  ➤ Require confirm ation of FDA approved, compendia supported, or Medicaid covered diagnosis  ➤ Click here for a copy of the CNS stimulant for patients 18 years and older fax form.  BOSE OPTIMIZATION (I/O)  ➤ See Dose Optimization Chart for affected drugs and strengths  FRE QUENCY (QUANTITY/DURATION (FQ/D))  ➤ Quantity limits based on daily dosage as determined by FDA labeling  ➤ Quantity limits for patients less than 18 years of age to include:  ■ Short-acting CNS stimulants not to exceed 3 dosage units daily with maximum of 90 days per strength (for titration)  ■ Long-acting CNS stimulants not to exceed 1 dosage unit daily with maximum of 90 days.  Concerta 36mg not to exceed 2 units daily.  ➤ Guantity limits for patients 18 years of age and older to include:  ■ Short-acting CNS stimulants: not to exceed 3 dosage units daily with maximum of 30 days.  Concerta 36mg not to exceed 2 units daily.  ➤ For patients 18 years of age and older: a 90 day supply may be obtained with confirm ation of FDA approved, Compendia supported or Medicaid covered diagnosis
	Multiple So	clerosis Agents
Avonex <sup>6</sup> Betaseron <sup>6</sup> Copaxone <sup>6</sup> 20 mg/mL Gilenya <sup>6 ≋T</sup>	Aubagio <sup>©ST</sup> Copaxone <sup>®</sup> 40 mg/mL Extavia <sup>®</sup> Glatopa™ Plegridy <sup>®</sup> Rebif <sup>®</sup> Tecfidera <sup>® ST</sup>	STEP THE RAPY (ST)  ➤ Gilenya™ (fingolimod) – requires a trial with a preferred injectable product  ➤ Aubagio® (teriflunomide) and Tecfidera™ (dimethyl fumarate) – require a trial with a preferred oral agent
		ine Receptor Agonists

1 = Preferred as of 11/19/2015

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<sup>2 =</sup> Non-preferred as of 11 / 19/2015

Preferro	ed Drugs	Non-Preferred Drugs		Prior Authorization/Coverage Parameters
pramipexole		Mirapex <sup>®</sup> Mirapex ER <sup>®</sup> Neupro <sup>®</sup> pramipexole ER	Requip <sup>©</sup> Requip <sup>®</sup> XL <sup>™00</sup> ropinirole ER	DOSE OPTIMIZATION (DO) ➤ See Dose Optimization Chart for affected strengths
	Other Agents for Attention Defi			cit Hyperactivity Disorder (ADHD) <sup>66</sup>
guan facine ER 00	Strattera <sup>© 00</sup>	clonidine ER Intuniv <sup>© 00</sup>	Kapvay <sup>®</sup>	CLINICAL CRITERIA (CC)  ➤ Confirm diagnosis for an FDA-approved or Compendia supported indication for beneficiaries < 18 years of age.  ➤ Prior authorization is required for initial prescriptions for non-stimulant therapy for beneficiaries less than 6 years of age  DOSE OPTIMIZATION (DO)  ➤ See Dose Optimization Chart for affected strengths

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	ypnotics/Sleep Agents	
estazolam es. Foro flurazepam es. Foro tema zepam 15mg, 30mg es. Foro zolpidem foro	Ambien GF 10/10  Ambien CR F 10/10  Belsomra F 10/10  Edluar F 10/10  eszopiclone F 10/10  Halcion G 20, F 10/10  Intermezzo G F 10/10  Lunesta G 10, F 10/10  Restorii G 20, F 10/10  Rozerem G F 10/10  Silenor G Sonata G 10/10  temazepam 7.5mg, 22.5mg G 10/10  triazolam G 10/10  zaleplon F 10/10  zolpidem ER f 10/10	DOSE OPTIMIZATION (DO)

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	Selective Seroto	nin Reuptake Inhibitors (SSRIs)
citalopram escitalopram escitalopram (tablet) fluoxetine 10mg, 20mg, 40mg fluoxetine (solution) paroxetine sertraline	Brintellix® paroxetine CR Brisdelle® Paxil® Celexa® Paxil CR® escitalopram Pexeva® (solution) Prozac® fluoxetine 60 mg fluoxetine DR weekly fluvoxamine ER® Lexapro® 00	DOSE OPTIMIZATION (DO)  ➤ See Dose Optimization Chart for affected strengths  CLINICAL CRITERIA (CC)  ➤ Clinical editing will allo wpatients currently stabilized on fluvoxamine or fluvoxamine ER to continue to receive that agent without PA  ➤ Clinical editing to allowpatients with a diagnosis of Obsessive Compulsive Disorder (OCD) to receive fluvoxamine and fluvoxamine ER without prior authorization
		ohrine Reuptake Inhibitors (SNRIs) <sup>ST</sup>
duloxetine venlafaxine venlafaxine ER (capsule)	Cymbalta <sup>©</sup> Desvenlafaxine base ER Desvenlafaxine fumarate ER E ffexor XR <sup>© 00</sup> Fetzima <sup>©</sup> Khedezla <sup>™</sup> P ristiq <sup>© 00</sup> Savella <sup>©</sup> venlafaxine ER (tablet)	DOSE OPTIMIZATION (DO)  ➤ See Dose Optimization Chart for affected strengths  STEP THE RAPY (ST)  ➤ Trial of an SSRI prior to an SNRI  • Step therapy is not required for the following indications:  • Chronic musculoskeletal pain (CMP)  • Diabetic peripheral neuropathy (DPN)  • Fibrom yalgia (FM)  ➤ Cym balta  (duloxetine) — Requires a trial with a tricyclic antidepressant OR gabapentin for treatment of Diabetic Peripheral Neuropathy (DPN)

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorizat	ion/Coverage Parameters
	Serotonin Re	ceptor Agonists (Triptans)	
rizatriptan F/2/0	almotriptan <sup>FO /D</sup>	FRE QUENCY/QUANTITY/DURATION (F/	Q.ID)
sumatriptan FIGTO	Amerge Fran	almotriptan	18 units every 30 days
	Axert <sup>® Frazo</sup> Frova <sup>® Frazo</sup>	Amerge <sup>●</sup>	
	Imitrex <sup>© FO/ID</sup>	Axert® 6.25mg	
	Maxatt <sup>e</sup> Fort	Frova <sup>®</sup>	
	naratriptan Froro	Im itrex <sup>®</sup> tablets	
	Relpax <sup>e ⊧∧</sup> /0	Imitrex <sup>®</sup> Nasal Spray	
	Sum avel <sup>®</sup> DosePro <sup>™</sup>	Naratriptan	
	Treximet <sup>® Forto</sup>	Relpax <sup>®</sup> 20mg	
	zolm itriptan	sumatriptan tablets	
	Zomig <sup>e F∧2/0</sup>	Treximet <sup>6</sup>	
		Sum avel <sup>®</sup> DosePro	
		zolmitriptan (tablet, ODT) 2.5mg	
		zolmitriptan (tablet, ODT) 5mg	
		Zomig/Zomig <sup>®</sup> ZMT 2.5mg	
		Zomig <sup>®</sup> /Zomig® ZMT 5mg	
		Zomig <sup>®</sup> Nasal Spray	
		Axert <sup>®</sup> 12.5mg	24 tablets every 30 days
		Maxalt Maxalt MLT®	
		Relpax <sup>®</sup> 40mg	
		rizatriptan (tablet, ODT)	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters		
V. DERMATOLOGIC AGENTS				
	Agents for Actinic Keratosis			
Aldara <sup>1</sup> diclofenac 3% gel <sup>Fraro</sup> Efudex <sup>6</sup> fluorouracil (solution) fluorouracil 0.5% cream (generic for Carac)	Carac <sup>®</sup> fluorouracil 5% cream (generic for Efudex cream) <sup>2</sup> imiquimod <sup>2</sup> Picato <sup>2</sup> Solaraze <sup>® FA270</sup> Tolak <sup>™</sup> Zydara <sup>2</sup>	FRE QUENCY/QUANTITY/DURATION (F/Q/D)  Solaraze / dictofenac 3% qet:  Maximum 100 (one hundred) grams as a 90 day supply Limited to one (1) prescription per year		
	Antibio	tics – Topical		
Altabax <sup>®</sup> mupirocin	Bactroban <sup>©</sup> Bactroban Nasal <sup>©</sup> (ointment) <sup>©©</sup> Centany <sup>©</sup> (ointment)	CLINICAL CRITERIA      Bactroban Nasal <sup>6</sup> ointment — Patient-specific considerations for drug selection include concerns related to use for the eradication of nasal colonization with methicillin-resistant Staphylococcus aureus (MRSA) in patients older than 12 years.		
	Anti-F un	gals – Topical		
dotrimazole OTC Lamisil AT <sup>®</sup> miconazole OTC Nyamyc™ nystatin (cream, ointment, powder) nystatin/triamcinolone Nystop <sup>®</sup> terbinafine OTC tolnaftate OTC	Alevazol OTC ST Cidodan (cream) ST cidopirox (cream, gel, suspension) ST clotrimazole / betamethasone ST clotrimazole Rx ST econazole ST Extraczo ST Exelderm ST Extina ST ketoconazole ST Luzu ST Mentax ST natifine ST Nation ST Oxistat ST Vusion ST	**STEP THERAPY (ST)  ➤ Trial of a preferred product (of comparable coverage) before using a non-preferred product FREQUENCY/QUANTITY/DURATION (F/Q/D)  ➤ Vusion® 50qm ointment — Maximum 100 (one hundred) grams in a 90 day time period		

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

#### NYS Medicaid Fee-For-Service Preferred Drug List

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
		tives, Topical
Benzadin pump	Acanya <sup>6</sup>	
dindam yein (gel, lotion, solution)	Benzadin <sup>e</sup> (gel)	
erythromycin (gel, solution)	Benzamycin <sup>€</sup>	
, , , , ,	Cleocin T	
	Clindadin <sup>©</sup>	
	Clindagel <sup>®</sup>	
	clindam ydin (foam , pledget)	
	clindamycin / benzoyl peroxide	
	Duac <sup>e</sup>	
	Erygel <sup>●</sup>	
	erythromycin (pledget)	
	erythromycin / benzoyl peroxide	
	E voclin <sup>®</sup>	
	Neuac <sup>®</sup>	
	Onexton™	
	Anti-Vir	als – Topical
Abreva <sup>6</sup>	acydovir (ointment)	
Zovirax <sup>●</sup> (cream.)	Denavir <sup>®</sup>	
	Sitavig <b>⁵</b>	
	Xerese <sup>●</sup>	
	Zovirax <sup>®</sup> (ointm ent)	
	lmmun omo du l	ators – Topical <sup>CDRP</sup>
Elidel <sup>6</sup> Protopic <sup>6</sup>	tacrolimus	CLINICAL DRUG REVIEW PROGRAM (CDRP)
		> All prescriptions require prior authorization
		> Refills on prescriptions are allowed
		➤ Click here for CDRP Topical Immunomodulators Prescriber Worksheet
	Psoriasis A	gents – Topical
calcipotriene (cream, ointment, scalp	calcipotriene / betamethasone dipropionate	
solution)	Calcitrene <sup>™</sup> (ointment)	
	calcitriol (ointment)	
	Dovonex <sup>e</sup> (cream)	
	Enstilar™	
	Sorilux <sup>®</sup>	
	Taclonex	
	Taclonex® Scalp®	
	Vectical <sup>®</sup>	

<sup>1 =</sup> Preferred as of 11/19/2015

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<sup>2 =</sup> Non-preferred as of 11 / 19/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters				
	Steroids, Topical – Low Potency					
hydrocortisone acetate OTC hydrocortisone acetate Rx hydrocortisone/ aloe vera OTC	alclometasone state fluocinolone (oil) state f	STEP THERAPY (ST)  Trial of preferred product (of comparable potency) before using non-preferred product.				
	Steroids, Topic	al – Medium Potency				
docortolone hydrocortisone butyrate (ointment, solution) hydrocortisone valerate mometasone furoate	Cloderm SI Cordran SI Cordran SI Cutivate SI Dermatop SI Elocon SI fluocinolone acetonide (cream, ointment, soln) SI fluticasone propionate SI hydrocortisone butyrate (cream) SI Luxiq SI Pandel SI prednicarbate SI Synalar SI	STEP THE RAPY (ST)  ➤ Trial of preferred product (of comparable potency) before using non-preferred product				
		cal – High Potency				
betamethasone dipropionate (cream, lotion) betamethasone valerate (cream, ointment) fluocinonide (cream, gel, solution) fluocinonide emollient fluocinonide-E triamcinolone acetonide	amcinonide <sup>ST</sup> Apexicon-E <sup>6ST</sup> betamethasone dipropionate (gel,ointment) <sup>ST</sup> betamethasone dipropionate, augmented <sup>ST</sup> betamethasone valerate (foam, lotion) <sup>ST</sup> desoximetasone <sup>ST</sup> difforasone <sup>ST</sup> difforasone <sup>ST</sup> Diprolene <sup>6</sup> ST Diprolene <sup>6</sup> AF <sup>ST</sup> fluocinonide 0.1% cream (generic for Vanos) <sup>ST</sup> fluocinonide (ointment) <sup>ST</sup> Halog <sup>6</sup> ST Kenalog <sup>6</sup> ST P sorcon <sup>ST</sup> Topicort <sup>6</sup> ST triamcinolone spray <sup>ST</sup> Trianex <sup>6</sup> ST Vanos <sup>6</sup> ST	STEP THE RAPY (ST)  ➤ Trial of preferred product (of comparable potency) before using non-preferred product				

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11 /1 9/2015

Prefe	erred Drugs	Non-Pre	ferred Drugs	Prior Authorization/Coverage Parameters
			Steroids, Topica	I – Very High Potency
dobetasol (cream , gel, ointment, solution) halobetasol		clobetasol (foam, lo Clobex <sup>©</sup> ST Olux <sup>©</sup> ST Olux-E <sup>©</sup> ST Temovate <sup>©</sup> ST Temovate-E <sup>©</sup> ST Ultravate	tion, spray) <sup>sr</sup>	STEP THERAPY (ST)  > Trial of preferred product (of comparable potency) before using non-preferred product.
			VI. ENDOCRINE AN	ID METABOLIC AGENTS
			Alpha-Gluco:	sidase Inhibitors <sup>8T</sup>
acarbose	Glyset <sup>®</sup>	Precose <sup>6</sup>		STEP THERAPY (ST)  Requires a trial with metformin with or without insulin prior to initiating alpha-glucosidase inhibitor therapy, unless there is a documented contraindication.
			Amylir	n Analogs <sup>8T</sup>
Symlin <sup>€</sup>		None		Requires a trial with metformin with or without insulin prior to initiating amylin analogue therapy, unless there is a documented contraindication.
			Anabolic Steroi	ds – Topical CORP FIGUD
Androgel <b>®</b>		Androderm <sup>©</sup> Axiron <sup>©</sup> Fortesta <sup>©</sup> Natesto™	Testim <sup>© Z</sup> testosterone gel Vogelxo <sup>©</sup>	CLINICAL DRUG RE VIEW PROGRAM (CDRP)  ➤ For diagnosis of hypogonadotropic or primary hypogonadism:  • Requires documented low testosterone concentration with two tests prior to initiation of therapy.  • Require documented testosterone therapeutic concentration to confirm response after initiation of therapy.  ➤ For diagnosis of delayed puberty:  • Requires documentation that growth hormone deficiency has been ruled out prior to initiation of therapy.  ➤ Click here for a copy of the Anabolic Steroid fax form  FRE QUENCY/QUANTITY/DURATION (F/Q/D)  ➤ Limitations for anabolic steroid products based on approved FDA labeled daily dosing and documented diagnosis:  Duration limit of six (6) months for delayed puberty

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Dru	ıgs	Non-Preferred Drugs		Prior Authorization/Coverage Pa	rameters
			Big	juanides	
metformin HCI metformin ER (generic for Glucophage XR*) Glucophage XR* Glucophage XR* Glumetza* metformin ER (generic for Fortamet*) Riomet*(solution)					
			Bisphospho	onates – Oral <sup>Floro</sup>	
alen dron ate		Actonel <sup>®</sup> Atelvia <sup>®</sup> Binosto <sup>®</sup> Boniva <sup>®</sup> Fosamax <sup>®</sup> Fosamax <sup>®</sup> Fosamax <sup>®</sup> Plus D Ibandronate risedronate		FRE QUENCY/QUANTITY/DURATION (F/Q/D)  Actonel 150 mg Boniva 150 mg ibandronate sodium 150 mg risedronate sodium 150 mg Actonel 35 mg alendronate sodium 35 mg alendronate sodium 70 mg Atelvia 35 mg Fosam ax 570 mg Fosam ax 70 mg Fosam ax 91 mg Fosam ax 91 mg Fosam ax 91 mg Fosam ax 92 mg Fosam ax 93 mg Fosam ax 94 mg Fosam ax 95 mg Fosam ax 95 mg Fosam ax 96 mg Fosam ax 970 mg	1 tablet every 28 days 4 tablets every 28 days 4 tablets every 28 days 4 bottles every 28 days
		'	Calcitoni	ns – Intranasal	
calcitonin-salmon Miac	calcin <sup>6</sup>	Fortical <sup>6</sup>			
			Dipeptidyl Peptidas	se-4 (DPP-4) Inhibitors <sup>ST</sup>	
Janumet <sup>©</sup> Jentad Janumet <sup>©</sup> XR Tradje Januvia <sup>© 10</sup>	dueto <sup>®</sup> enta <sup>®</sup>	Glyxam bi <sup>©</sup> Kazano™ Kom biglyze XR <sup>™</sup>	Nesina™ Onglyza <sup>© 00</sup> Oseni™	DOSE OPTIMIZATION (DO)  ➤ See Dose Optimization Chart for affected strengths  STEP THE RAPY (ST)  ➤ Requires a trial with metformin with or without insulin prior there is a documented contraindication.	to DPP-4 Inhibitor therapy, unles
			Glucagon-like Pepti	ide-1 (GLP-1) Agonists <sup>ST</sup>	
Bydureon <sup>®</sup> Tanze Byetta <sup>®</sup>	eum <sup>©</sup>	Trulicity™ Victoza <sup>®</sup>		STEP THERAPY (ST)   Requires a trial with metformin plus another oral antidiabet   Prior authorization is required with lack of covered diagnos	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 / 19/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
	Glucocor	ticoids – Oral
cortisone dexamethasone (tablet, solution) hydrocortisone methylprednisolone (4mg, 8mg, 32mg) methylprednisolone (dose-pack) prednisolone (solution) prednisone (dose-pack, solution, tablet)	budesonide EC Cortef <sup>®</sup> dexamethasone (elixir) dexamethasone intensol Dexpak <sup>®</sup> Entocort EC <sup>®</sup> Medrol <sup>®</sup> (dose-pack, tablet) methylprednisolone 16mg Millipred <sup>®</sup> Orapred <sup>®</sup> prednisolone ODT prednisone intensol Rayos <sup>®</sup> Uceris <sup>®</sup>	
	Growth Ho	rmones <sup>CC</sup> CORP
Genotropin	Humatrope <sup>®</sup> Tev-Tropin <sup>®</sup> Omnitrope <sup>®</sup> Zomacton <sup>®</sup> Saizen <sup>®</sup> Zorbtive <sup>®</sup>	CLINICAL DRUG RE VIEW PROGRAM (CDRP)  Prescriptions for enrollees that are 21 years of age or older require P A under the CDRP  Prescribers, not authorized agents, are required to call the clinical call center toll free number 1-877-309-9493 and respond to a series of questions that identify prescriber, patient and reason for prescribing a drug in this class for enrollees 21 years of age or older  Refills on prescriptions are allowed  Refer to the Preferred Drug Program web page and review list of preferred and non-preferred drugs when prescribing for enrollees under the age of 21  Click here for a copy of the CDRP Growth Hormone Prescriber Fax Form and Instructions CLINICAL CRITERIA (CC)  Patient-specific considerations for drug selection include concerns related to use of a non-preferred agent for FDA approved indications that are not listed for a preferred agent.  Appropriate diagnosis is required for all Growth Hormones, regardless of age or preferred status.
	Insulin –	Long-Acting
Lantus <sup>®</sup> Levemir <sup>®</sup>	Toujeo <sup>®</sup> Tresiba <sup>®</sup>	
	Insuli	n – Mixes
Humalog <sup>®</sup> Mix Novolog <sup>®</sup> Mix	None	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Prefer	red Drugs	Non-Prefe	erred Drugs	Prior Authorization/Coverage Parameters
			Insulin –	Rapid-Acting
Apidra <sup>●</sup> Humalog <sup>●</sup> 100 U <i>l</i> mL	Novolog <sup>©</sup>	Afrezza <sup>€</sup> Humalog <sup>€</sup> 200 U/m L <sup>2</sup>		
			Megl	itinides <sup>ST</sup>
nateglinide	repaglinide	Prandim et <sup>©</sup> Prandin <sup>©</sup>	repaglinide/met formin Starlix <sup>6</sup>	STEP THERAPY (ST)  Requires a trial with metformin with or without insulin prior to initiating meglitinide therapy, unless there is a documented contraindication.
	•		Pancrea	atic Enzymes
Creon <sup>6</sup> pancrelipase	Zenpep <sup>6</sup>	P ancrea ze <sup>€</sup> P ertzye <sup>™</sup>	Ultresa <sup>6</sup> Viokace <sup>6</sup>	
		Sodio	ım Glucose Co-Tran	sporter 2 (SGLT2) Inhibitors <sup>ST</sup>
Invokana <sup>6</sup>		Farxiga™ Invokamet <sup>6</sup> Jardiance <sup>6</sup>	Synjardy <sup>6</sup> Xigduo™ XR	STEP THERAPY (ST) ➤ Requires a trial with metformin with or without insulin prior to initiating SGLT2 Inhibitor therapy, unless there is a documented contraindication.
		•	Thiazolidine	ediones (TZDs) <sup>ST</sup>
pioglitazone		Actoplus Met <sup>®</sup> Actoplus Met <sup>®</sup> XR <sup>00</sup> Actos <sup>© 00</sup> Avandamet <sup>©</sup> Avandia <sup>©</sup> Duetact <sup>©</sup> pioglitazone / glim epi		DOSE OPTIMIZATION (DO)  ➤ See Dose Optimization Chart for affected strengths  STEP THERAPY (ST)  ➤ Requires a trial with metformin with or without insulin prior to initiating TZD therapy, unless there is a documented contraindication.

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11/19/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
	VII. GASTROINTESTINAL				
	Anti	i-Emetics			
ondansetron (ODT, solution, tablet)	Anzemet <sup>©</sup> granisetron (tablet) Sancuso <sup>©</sup> Zofran <sup>©</sup> (ODT, solution, tablet) Zuplenz <sup>©</sup>				
	G astrointes	stinal Antibiotics			
metronidazole (tablet) neomycin vancomycin	Ainia® Dificid® Flagyl® Flagyl® ER metronidazole (capsule) paromornycin Tindamax® tinidazole Vancocin® Xi faxan® 22.87.5070	CUNICAL CRITERIA (CC)  ➤ Xifaxan  — Requires confirmation of diagnosis of Traveler's diarrhea, hepatic encephalopathy, or irritable bowel syndrome with diarrhea (IBS-D)  STEP THE RAPY (ST)  ➤ Xifaxan  — Requires trial of a preferred fluoroquinolone antibiotic before rifaximin for treatment of Traveler's diarrhea  QUANTITY LIMITS:  ➤ Xifaxan:  ■ Traveler's diarrhea (200 mg tablet) — 9 (nine) tablets per 30 days (Dose = 200 mg three times a day for three days)  ■ Hepatic encephalopathy (550 mg tablets) — 60 tablets per 30 days (Dose = 550 mg twice a day)  ■ Irritable bowel syndrome with diarrhea (550 mg tablets) — 42 tablets per 30 days (Dose = 550 mg three times a day for 14 days)  ■ Maximum of 42 days supply (126 units) per 365 (three rounds of therapy).			

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
	Gastrointestinal Preparatory Agents				
Clearlax <sup>6</sup>	Colyte <sup>®</sup>				
Gavilax <sup>6</sup>	Gavilyte <sup>6</sup> -N				
Gavilyte <sup>®</sup> -C	Golytely <sup>®</sup>				
Gavilyte <sup>6</sup> -G	Moviprep <sup>6</sup>				
Glycolax <sup>9</sup>	Nulytely <sup>6</sup>				
Miralax <sup>®</sup> OTC	Osmoprep <sup>●</sup>				
PEG 3350 powder OTC	PEG 3350 powder pack OTC				
PEG 3350 / electrolytes solution Rx	PEG 3350 with flavor packs				
	Prepopik <sup>®</sup>				
	Suprep <sup>6</sup>				
	Trilyte <sup>©</sup>				
Helicobacter pylori Agents					
lansoprazole / am oxicillin / clarithromycin	Omeclamox-Pak <sup>®</sup>				
Pylera <sup>€</sup>	Prevpac <sup>●</sup>				

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 / 19/2015

Preferre	ed Drugs	Non-Preferr	ed Drugs	Prior Authorization/Coverage Parameters
Proton Purr				nhibitors (PPIs) FIGID
omeprazole Rx pantoprazole Prilosec <sup>©</sup> OTC		Aciphex <sup>®</sup> Dexilant ™00 esomeprazole magnesiu Nexium) Esomeprazole Strontium lansoprazole Rx (capsul Nexium ®RX 00 omeprazole OTC omeprazole/sodium bica Prevacid OTC Prevacid ®Rx 00 Prilosec ®Rx Protonix 00 Zegerid 00	e,ODT)	DOSE OPTIMIZATION (DO)  ➤ See Dose Optimization Chart for affected strengths  FRE QUENCY/QUANTITY/DURATION (F/Q/D)  ➤ Quantity limits:  • Once daily dosing (30 units every 30 days) for:  • GERD  • erosive esophagitis  • healing and maintenance of duodenal/gastric ulcers (including NSAID-induced)  • prevention of NSAID-induced ulcers  • Twice daily dosing (60 units every 30 days) for:  • hypersecretory conditions  • Barrett's esophagitis  • H. pylori  • refractory GERD  ➤ Duration limits:  • 60 days for:  • Mild/moderate GERD  • acute healing of duodenal/gastric ulcers (including NSAID-induced)  • 365 days for:  • Maintenance treatment of duodenal ulcers  • 14 days for:  • H. pylori
			Sulfasalaz	rine Derivatives
Apriso <sup>©</sup> Delzicol™ Dipentum <sup>©</sup> sulfasalazine DRÆC	sulfasalazine IR sulfazine sulfazine EC	Asacol HD <sup>©</sup> Azul fidine <sup>©</sup> Azul fidine Entab <sup>©</sup> bal salazide	Colazal <sup>©</sup> Giazo <sup>™</sup> Lialda <sup>©</sup> Pentasa <sup>©</sup>	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferr	ed Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters
		VIII. HEMA	TOLOGICAL AGENTS
		Anticoa	gulants – Injectable
enoxaparin sodium	Fragmin <sup>6</sup>	Arixtra <sup>© 22</sup> Lovenox <sup>©</sup> fondaparinux <sup>22</sup>	CLINICAL CRITERIA (CC)      Clinical editing to allo wip atients with a diagnosis of Heparin Induced Thrombocytopenia (HIT) to receive fondaparinux (Arixtra®) without prior authorization.
		Antic	o agulants – Oral
Courn adin <sup>6</sup> Jantoven <sup>6</sup> Prada xa <sup>6</sup>	warfarin Xarelto <sup>©</sup>	Eliquis <sup>e <u>se</u> Savaysa<sup>e</sup> Xarelto<sup>e</sup> (dose pack)</sup>	CLINICAL CRITERIA (CC)     Clinical editing will allow patients currently stabilized on Eliquis® (apixaban) to continue to receive that agent without PA
	•	Erythropoiesis S	Stimulating Agents (ESAs) <sup>CC</sup>
Aranesp <sup>®</sup>	Procrit <sup>®</sup>	Epogen <sup>®</sup> Mircera <sup>®</sup>	CUNICAL CRITERIA (CC)  > Confirm diagnosis for FDA or Compendia supported uses
		Pla	telet Inhibitors
Aggrenox <sup>®</sup> Brilinta <sup>®</sup> clopidogrel	dipyridamole Effient <sup>©</sup>	dipyridamole / aspirin Durla za <sup>©</sup> Persantine <sup>©</sup> Plavix <sup>©</sup> tid opidine Zontivity <sup>©</sup>	
		IX. IMMU	INOLOGIC AGENTS
		lmmunomod	ulators – Systemic <sup>CC, 8T</sup>
Enbrel <sup>©</sup>	Humira <sup>©</sup>	Actem ra <sup>®</sup> (subcutaneous) Cim zia <sup>®</sup> Cosentyx <sup>®</sup> Kineret <sup>®</sup> Orencia <sup>®</sup> (subcutaneous) Otezia <sup>®</sup> Simponi <sup>®</sup> Stelara <sup>®</sup> Xelianz <sup>®</sup>	CLINICAL CRITERIA (CC) Confirm diagnosis for FDA or Compendia supported uses  STEP THERAPY (ST) Trial of a disease-modifying anti-rheumatic drug (DMARD) prior to treatment with an immunomodulator

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
X. MISCELLANEOUS AGENTS					
Progestins (for Cachexia)					
megestrol acetate (suspension)	Megace <sup>®</sup> (suspension) Megace ES <sup>®</sup> megestrol ES (suspension)				
	Epinephrin	e, Self-injected			
Adrenaclick EpiPen Auvi- Q <sup>®</sup> EpiPen Jr epinephrine	None				
	XI. MUSCULOS	KELETAL AGENTS			
	Skeletal Mu	iscle Relaxants			
chlorzoxazone cyclobenzaprine 5m g, 10mg dantrolene methocarbamol orphenadrine ER tizanidine (tablet)	metaxalone Parafon Forte <sup>®</sup> DSC Robaxin <sup>®</sup> Skelaxin <sup>®</sup> Soma <sup>® ST, Frano Soma<sup>®</sup> 250 <sup>ST, Frano</sup></sup>	CLINICAL CRITERIA (CC)  For carisoprodol/codeine products:  Limited to a total of four (4) opioid prescriptions every 30 days; Exemption for diagnosis of cancer or sickle cell disease.  Medical necessity rationale for opioid therapy is required for patients on established buprenorphine therapy.  P A required for initiation of opioid therapy in patients currently on benzodiazepine therapy.  STEP THE RAPY (ST)  Trial with one (1) preferred analgesic and two (2) preferred skeletal muscle relaxants prior to use of carisoprodol containing products:  carisoprodol  carisoprodol/AS A  carisoprodol/AS A/codeine  Som a  FRE QUENCY/QUANTITY/DURATION (F/Q/D)  Maximum 84 cumulative units per a year  Carisoprodol — Maximum 4 (four) units per day, 21 day supply  Carisoprodol combinations — Maximum 8 (eight) units per day, 21 (twenty-one) day supply (not to exceed the 84 cumulative units per year limit)			

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
XII. OPHTHALMICS					
	Alpha-2 Adrenergic Agonists (for Glaucoma) – Ophthalmic				
Alphagan P <sup>●</sup> Simbrinza™	apracionidine lopidine				
brim onidine 0.2%	brim onidine 0.15%				
	Antibiotics	s – Ophthalmic			
bacitracin / polymyxin B	Azasite <sup>6</sup>				
erythrom ycin	bacitracin				
gentamicin	Bleph <sup>6</sup> -10				
llotycin™	Garamycin <sup>●</sup>				
Natacyn <sup>6</sup>	neomycin / bacitracin / polymyxin				
neom ycin / gram icidin / polym yxin	Neosporin <sup>6</sup>				
polym yxin / trim ethoprim	P olytrim <sup>6</sup>				
sulfacetamide (solution)	sulfacetamide (ointment)				
tobramydin	Tobrex <sup>®</sup>				
	Antibiotics/Steroid Co	ombinations – Ophthalmic			
Blephamide <sup>6</sup>	Maxitrol <sup>®</sup>				
neomycin/polymyxin/dexamethasone	neomycin / bacitracin / polymyxin / HCTZ				
sulfacetamide / prednisolone	neomycin / polymyxin / HCTZ				
TobraDex <sup>®</sup> (ointment, suspension)	Pred-G <sup>●</sup>				
	TobraDex <sup>®</sup> ST				
	tobramycin / dexametha sone (suspension)				
	Zylet <sup>●</sup>				
	Antihistamines – Ophthalmic				
Pataday <sup>6</sup>	azelastine Lastacaft <sup>e</sup>				
	Bepre ve <sup>®</sup> Optivar <sup>®</sup>				
	Elestat <sup>6</sup> Patanol <sup>6</sup>				
	Em adine <sup>6</sup> Pazeo™				
	epinastine				

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Pre	ferred Drugs	Non-Pi	eferred Drugs	Prior Authorization/Coverage Parameters
			Beta Bl	ockers – Ophthalmic
betaxolol Betimol® Betoptic S® carteolol Combigan® Istalol® levobunolol metipranolol timolol maleate (gel, solution)		Betagan <sup>®</sup> Tim optic <sup>®</sup> Tim optic <sup>®</sup> in Ocud Tim optic-XE <sup>®</sup>	ose <sup>6</sup>	
			Fluoroquir	nolones – Ophthalmic <sup>8T</sup>
ciprofloxacin Vigamox <sup>®</sup> Besivance <sup>®</sup> Moxeza <sup>®</sup> ofloxacin Ciloxan <sup>®</sup> Ocuflox <sup>®</sup> gatifloxacin Zymaxid <sup>®</sup> levofloxacin		Ocuflox <sup>®</sup>	STEP THERAPY (ST)  ➤ For patients 21 years or younger, attempt treatment with a non-fluoroquinolone ophthalmic antibiotic before progressing to the a fluoroquinolone ophthalmic product	
	<u> </u>	Non-S	teroidal Anti-Inflam	nmatory Drugs (NSAIDS) – Ophthalmic
didofenac ketorolac Acular <sup>®</sup> Ilevro <sup>®</sup> flurbiprofen Acular LS <sup>®</sup> Nevanac <sup>®</sup> Acuvail <sup>®</sup> Ocufen <sup>®</sup> bromfenac Prolensa™		Nevanac <sup>6</sup> Ocufen <sup>6</sup>		
		-	Prostagland	in Agonists – Ophthalmic
latanoprost		birnatoprost Lumigan <sup>s</sup> Travatan Z <sup>s</sup>	travoprost Xalatan <sup>e</sup> Zioptan™	

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs		referred Drugs	Prior Authorization/Coverage Parameters					
XIII. OTICS								
		Fluoroquii	nolones – Otic					
Ciprodex <sup>6</sup> offoxacin Cipro HC <sup>6</sup>								
		XIV. RENAL AN	D GENITOURINARY					
		Alpha Reductas	e Inhibitors for BPH					
	Avodart <sup>e</sup>							
		ulosin						
	·							
		Cystine Den	leting Agents <sup>©</sup>					
	Procysbi <sup>6 ST</sup>	-,	CLINICAL CRITERIA (CC)					
			Confirm diagnosis of nephropathic cystinosis					
			STEP THERAPY (ST)					
			> Requires a trial with Cystagon immediate-release capsules					
		Dhoenhato Ri	1 - 1					
	1							
Renagel								
	Phoslyra	·						
		Selective Alpha	Adrenergic Blockers					
tamsulosin	Flomax	Uroxatral <sup>€</sup>						
	Rapa flo <sup>®</sup>							
		Urinary Tract	Antispasmodics					
Vesicare <sup>€ 00</sup>	Detrol <sup>®</sup>	oxybutynin ER <sup>00</sup>	DOSE OPTIMIZATION (DO)					
	Detrol LA <sup>● №</sup>	Oxytrol <sup>®</sup>	➤ See Dose Optimization Chart for affected strengths					
	Ditropan XL <sup>®</sup>	tolterodine						
	Enable x <sup>6 00</sup>	tolterodine ER						
	Gelnique <sup>®</sup>	trospium						
	Myrbetriq <sup>e</sup>	trospium ER						
	•	Xanthine Ox	idase Inhibitors					
	Uloric <sup>€</sup>	Zyloprim <sup>®</sup>						
	ofloxacin  ofloxacin  Fosrenol  Renagel  tam sulosin	offoxacin  Cipro HC®  Avodart® dut asteride dut asteride / tams Jalyn® Proscar®  Procysbi®®™  Renagel®  Auryxia™ Phoslo® Phoslyra®  tamsulosin  Flomax Rapa flo®  Vesicare®®  Detrol® Detrol LA®® Detrol LA®® Celnique® Myrbetriq®	Fluoroquii  ofloxacin  Cipro HC®  XIV. RENAL AN  Alpha Reductas  Avodart® dut asteride dut asteride / tamsulosin Jalyn® Proscar®  Procysbi®®®T  Procysbi®®®T  Procysbi®®®T  Phosphate Bi  Fosrenol® Renagel® Phoslo® Phoslo® sevelam er carbonate Phoslyra® Velphoro®  Selective Alpha  tamsulosin  Flomax Rapa flo®  Uroxatral®  Vesicare®®®  Detrol® Detrol LA®®® Oxytrol® Ditrop an XL® totterodine E nablex®®® Totterodine ER Gelnique® trospium Myrbetriq® Txanthine Ox					

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs Non-Preferred Drugs			Prior Authorization/Coverage Parameters					
XV. RESPIRATORY								
Anticholinergics / COPD Agents								
Atrovent HF A <sup>®</sup> Combivent Respirat <sup>®</sup> ipratropium ipratropium / albuterol Spiriva <sup>®</sup>	Anoro Ellipta <sup>©</sup> Daliresp <sup>©</sup> Incruse Ellipta <sup>©</sup> Seebri Neohaler <sup>©</sup>	Spiriva Respimat <sup>©</sup> Stiolto Respimat <sup>©</sup> Tudorza Pressair <sup>©</sup> Utibron Neohaler <sup>©</sup>						
		Antihistami	ines – Intranasal					
Astepro™ Patanase <sup>6</sup>	azelastine	olopatadine						
	•	Antihistamines	- Second Generation					
cetirizine OTC (tablet) cetirizine OTC (syrup/solution 1mg/1mL) fexofenadine OTC (suspension) levocetirizine (tablet) loratadine OTC			CLINICAL CRITERIA (CC)  ➤ No prior authorization required for patients less than 24 months of age					

<sup>1 =</sup> Preferred as of 11/19/2015 2 = Non-preferred as of 11/19/2015

Pro	eferred Drugs		erred Drugs	Prior Authorization/Coverage Parameters		
		Beta	Adrenergic Agents	s – Inhaled Long-Acting 🥰 ROYD		
Brovana <sup>6</sup>	Serevent Diskus <sup>®</sup>	Arcapta <sup>™</sup>	Striverdi Respimat <sup>e</sup>	CLINICAL CRITERIA (CC)		
Foradil <sup>e</sup>		Perforomist <sup>e</sup>			-acting beta agonist prescriptions for beneficiaries under FDA o ndicated:	
					Arcapta™	≥18 years
					Brovana <sup>6</sup>	≥18 years
					Foradil <sup>®</sup>	≥ 5 years
					Perforomist <sup>€</sup>	≥18 years
					Serevent <sup>e</sup>	≥4 years
					Striverdi <sup>e</sup>	≥18 years
				FREQ	UENCY/QUANTITY/DUI	RATION (F/Q/D)
		Maximum units per 30 days				
					Arcapta <sup>™</sup>	30 units (1 box of 30 unit dose capsules)
					Brovana <sup>€</sup>	60 units (1 carton of 60 vials or 120 mL)
					Foradil <sup>e</sup>	60 units (1 box of 60 unit dose capsules)
					Perforomist <sup>e</sup>	60 units (1 carton of 60 vials or 120 mL)
					Serevent <sup>e</sup>	1 diskus (60 blisters)
					Striverdi <sup>e</sup>	1 unit (one cartridge and one Respirnat inhaler)
		В	eta₂ Adrenergic Age	nts –	Inhaled Short-Acting	
albuterol	Proventil HFA <sup>●</sup>	levalbuterol (solution)			_	
ProAir HFA <sup>®</sup>		Proair <sup>®</sup> RespiClick Ventolin HFA <sup>®</sup>	XopenexHFA <sup>●</sup>			

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 / 19/2015

Preferred Drugs	Non-Preferred Drugs	Prior Authorization/Coverage Parameters			
	Corticos	teroids – Inhaled <sup>FIQ/D</sup>			
Asmanex <sup>6</sup>	Aerospan <sup>6</sup>	FRE QUENCY/QUANTITY/DURATION (F/Q/D)			
Flovent Diskus <sup>6</sup>	Alvesco <sup>6</sup>	Aerospan <sup>®</sup> 80 mog	2 inhalers every 30 days		
Flovent HFA <sup>6</sup>	Arnuity Ellipta™	Alvesco® 80 m cg	1 inhaler every 30 days		
Pulmicont <sup>®</sup> Flexhaler	Asmanex <sup>®</sup> HFA	Alvesco® 160 mcg	1 inhaler every 30 days. Up to 1 inhaler every 15 days with previous or al corticosteroid use.		
QVAR <sup>®</sup>		Arnuity Ellipta	1 inhaler every 30 days		
		Asmanex 110 mcg	1 inhaler every 30 days		
		Asmanex 220 mcg (30 units)	1 inhaler every 30 days		
		Asman ex <sup>®</sup> 220 m og (80 units)	1 inhaler every 30 days Up to 1 inhaler every 15 days with previous oral corticosteroid use.		
		Asman ex = 220 m cg (120 units)	1 inhaler every 60 days. Up to 1 inhaler every 30 days with previous oral corticosteroid use.		
		Asmanex <sup>®</sup> HFA 100 mcg	1 inhaler every 30 days		
		Asmanex HFA 200 mcg	1 inhaler every 30 days		
		Flovent Diskus 50mcg, 100 mcg	1 diskus every 30 days		
		Flovent Diskus® 250 mog	1 diskus every 15 days. Up to 1 diskus every 7 days with previous or al corticosteroid use.		
		Flovent HFA 44mcg, 110 mcg	1 inhaler every 30 days		
		Flovent HFA <sup>®</sup> 220mcg	1 inhaler every 30 days. Up to 1 inhaler every 15 days with previous oral corticosteroid use.		
		Pulmicort90mcg	1 inhaler every 30 days		
		Pulmicort 180mcg	1 inhaler every 15 days		
		QVAR 40mcg	1 inhaler every 25 days		
		QVAR 80mcg	1 inhaler every 12 days		
	Corticosteroid/Beta₂ Adrenergic Age	ent (Long-Acting) Combinations – I	nhaled <sup>으େ ନଦାପ</sup>		
Advair Diskus ** Dulera **	Breo Ellipta <sup>®</sup>	CLINICAL CRITERIA (CC)			
AdvairHFA <sup>●</sup> Symibicont <sup>●</sup>		<ul> <li>PA is required for all newlong-a or compendia supported age as</li> </ul>	cting beta agonist prescriptions for beneficiaries under Fl indicated:		
		Advair Diskus	≥4 years		
		Advair HFA	≥12 years		
		Breo Ellipta™	≥18 years		
		Dulera -	≥12 years		
		Symbicort*	≥12 years		
		•	<del>'</del>		
		FRE QUENCY/QUANTITY/DURATIO			
		Advair Diskus	One (1) inhaler/diskus every 30 days		
		Advair HFA	<b>↓</b>		
		Breo Ellipta™	<b>↓</b>		
		Dulera <b>"</b>	<b>」</b>		
		Symbicort <sup>®</sup>			

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

Preferred Drugs	Non-Pro	eferred Drugs		r Authorization/Coverage Parameters			
Cortico steroids – Intranasal							
fluticasone	Beconase AQ€	QNASL*	FRE QUENCY/QUANTITY/DI	URATION (F/Q/D)			
Nasonex <sup>6</sup>	budesonide Dymista <sup>6</sup>	Rhinocort Aqua <sup>e</sup> triamcinolone	Beconase AQ <sup>●</sup>	One (1) inhaler every 22 days			
	Flonase <sup>6</sup>	Veramyst <sup>e</sup>	flunisolide	One (1) inhaler every 25 days			
	Flunisolide	Zetonna <sup>6</sup>	budesonide	One (1) inhaler every 30 days			
	Omnaris <sup>®</sup>		Dymista™				
			Flonase				
			fluticasone				
			Nasacort AQ <sup>●</sup>				
			Nasonex <sup>6</sup>				
			Om naris <sup>●</sup>				
		QNASL <sup>®</sup>					
			Rhinocort Aqua <sup>€</sup>				
			triamcinolone				
			Veram yst <sup>€</sup>				
			Zetonna™				
		Leukotr	iene Modifiers				
montelukast <sup>ਭਾ</sup>	Accolate <sup>6</sup>		STEP THE RAPY (ST)				
zafirlukast	Singulair <sup>e sr</sup>		<ul> <li>For non-asthmatic patients, trial of intranasal corticosteroid or a 2nd generation oral antihistamine before montelukast (Singulair<sup>6</sup>)</li> </ul>				

<sup>1 =</sup> Preferred as of 11/19/2015

<sup>2 =</sup> Non-preferred as of 11 /1 9/2015

#### NYS Medicaid Fee-For-Service Clinical Drug Review Program (CDRP)

The Clinical Drug Review Program (CDRP) is aimed at ensuring specific drugs are utilized in a medically appropriate manner.

Under the CDRP, certain drugs require prior authorization because there may be specific safety issues, public health concerns, the potential for fraud and abuse or the potential for significant overuse and misuse.

#### Prior Authorization

Prior authorization for some drugs subject to the CDRP must be obtained through a representative at the clinical call center. Prior authorization is required for original prescriptions, not refills. For some drugs subject to the CDRP, only prescribers, not their authorized agents, can initiate the prior authorization process.

Fax requests for prior authorization are not permitted. Each CDRP drug has specific clinical information that must be provided to the clinical call center before prior authorization will be issued. Prescribers may be asked to fax that information. Clinical guidelines for the CDRP as well as prior authorization worksheets are available online at <a href="http://newyork.fhsc.com/providers/CDRP">http://newyork.fhsc.com/providers/CDRP</a> forms, asp.

The following drugs are subject to the Clinical Drug Review Program:

- becaplermin gel (Regranex<sup>®</sup>)
- emtricitabine/tenofovir (Truvada<sup>®</sup>)
- fentanyl mucosal agents
- <u>lidocaine patch (Lidoderm<sup>®</sup>)</u>

- oxazolidinone antibiotics (Sivextro<sup>™</sup>, Zyvox<sup>®</sup>)
- palivizumab (Synagis<sup>®</sup>)
- sodium oxybate (Xyrem<sup>®</sup>)
- somatropin (Serostim<sup>®</sup>)

The following drug classes are subject to the Clinical Drug Review Program and are also included on the Preferred Drug List:

- Anabolic Steroids
- . Central Nervous System (CNS) Stimulants for 18 years and older
- Growth Hormones for 21 years and older
- Phosphodiesterase type-5 (PDE-5) Inhibitors for PAH
- Topical Immunomodulators

#### NYS Medicaid Fee-For-Service Drug Utilization Review (DUR) Program

Frequency/Quantity/Duration (F/Q/D) Program and Step Therapy parameters are implemented to ensure clinically appropriate and cost effective use of these drugs and drug classes.

For additional Step Therapy and Frequency/Quantity/Duration parameters for drugs and drug classes that are also included on the Preferred Drug List (PDL), please see pages 3 through 31.

Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Qua (F/Q/D) Par		Additional / Alternate Parameter(s)
Acthar <sup>6</sup> (ACTH injectable)	Requires trial of first-line therapy for all FDA- approved indications, other than infantile spasms.  Note: Acthar is first line therapy for infantile spasms in children less than 2 years of age — step therapy not required.  DURATION LIMITS: Infantile spasms — 3  Multiple sclerosis — 3  Puration Limits: Infantile spasms — 4  < 2 years of age  Multiple sclerosis — 3  Rheum atic disorders  Dermatologic condit  Allergic states (seru weeks		35 m L (seven 5 m L weeks; indicated for 5 weeks s – 5 weeks ions – 5 weeks	Confirm diagnosis for Medicaid covered uses. Medicaid Fee-For-Service benefit does not cover for diagnostic purposes.
	FDA Indication		First line Therapy	
	Multiple Scierosis (MS) exacerbations		Corticosteroid or plasmapheresis	
	Polym yositis/demiatom yositis		Corticosteroid	
	Idiopathic nephrotic syndrome		ACE Inhibitor, diuretic, corticosteroid (and for refractory patients: an immunosuppressive)	
	Systemic lupus erythematosus (SLE)		Corticosteroid, antimalarial, or cytotoxic/im munosuppressive agent	
	Nephrotic syndrome due to SLE		lm mum osuppressive	, corticosteroid, or ACE Inhibitor
	Rheumatic disorders (specifically: psoriatic ar arthritis, juvenile rheumatoid arthritis, ankylos	•	Corticosteroid, topical retinoid, biologic disease-modifying antirheumatic drugs (DMARD), non-biologic DMARD, or a non-steroidal anti-inflammatory drug (NSAID)	
	Dermatologic diseases (specifically Stevens-J erythema multiforme)	Johnson syndrome and	Corticosteroid or analgesic	
	Allergic states (specifically serum sickness)		Topical or oral corticosteroid, antihistamine, or NSAID	
	Ophthalmic diseases (keratitis, iritis, iridocydi uveitis/choroiditis, optic neuritis, chorioretinitis inflammation)		Analgesic, anti-infective agent, and agents to reduce inflammation, such as NSAIDs and steroids	
	Respiratory diseases (systemic sarcoidosis)		Oral costicosteroid or an imunosuppressive.	

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Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Amoxicillin ER (Moxatag <sup>®</sup> )	Prescribers should attempt treatment with a more cost effective immediate-release amoxicillin first before progressing to extended-release amoxicillin	QUANTITY LIMIT: > Equal to 10 tablets per fill	
Anabolic Steroids — Injectable  Depo-Testosterone  testosterone cypionate  testosterone enanthate  Anabolic Steroids — Oral  Anadrol-50  Android  Androxy <sup>TM</sup> Methitest  Oxandrin  oxandrolone  Testred		Limitations for anabolic steroid products is based on approved FDA labeled daily dosing and documented diagnosis not to exceed a 90-day supply (30-day supply for oxandrolone):  Initial duration limit of 3 months (for all products except oxandrolone), requiring documented follow-up monitoring for response and/or adverse effects before continuing treatment  Duration limit of 6 months for delayed puberty  Duration limit of 1 month for all uses of oxandrolone products	
Anti-Retroviral (ARV) Interventions		QUANTITY LIMITS:  > Limit ARV active ingredient duplication > Limit ARV utilization to a maximum of five products concurrently - excluding boosting with ritonavir (dose limit 600 mg or less) or cobicistat > Limit Protease Inhibitor utilization to a maximum of two products concurrently > Limit Integrase inhibitor utilization to a maximum of one product concurrently	Require confirmation of FDA approved or compendia supported use     Point of service edit for contraindicated antiretroviral / non-antiretroviral combinations     Point of service edit for contraindicated antiretroviral / antiretroviral combinations
Antidiabetic agents (not on the PDL)  chlorpropamide  glimepiride  glipizide (Glucotrol <sup>®</sup> , Glucotrol XL <sup>®</sup> )  glyburide (Diabeta <sup>®</sup> , Glynase <sup>®</sup> )  glyburide, micronized  tolazamide  tolbutamide	<ul> <li>Requires a trial with metformin with or without insulin prior to initiating other antidiabetic agents, unless there is a documented contraindication.</li> <li>Clinical editing to allow patients with a diagnosis of gestational diabetes to receive glyburide without a trial of metformin first.</li> </ul>		

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Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Antifungals, Topical — for Onychomycosis ➤ cidopirox 8% solution ➤ Jublia <sup>©</sup> ➤ Kerydin <sup>©</sup> ➤ Penlac <sup>©</sup>	Trial with an oral antifungal agent* prior to use of cidopirox 8% solution (P enlac)  * terbinafine (Lamisil®) tablets; griseofulvin (Grifulvin V®, Gris PEG®) oral suspension, ultramicronized tablets micronized tablets; itraconazole (Sporanox®, Onmel™) tablets, oral solution  Trial with ciclopirox 8% solution prior to the use of other topical antifungals [efinaconazole (Jublia) or tavaborole (Kerydin)]		
Becaplermin (Regranex <sup>®</sup> )		QUANTITY LIMIT: > 2 (two) 15 gram tubes in a lifetime	
Benzodiazepine agents – oral  > alprazolam (Niravam ™, Xanax Nanax XR)  > chlordiazepoxide (Librium Nanax N	<ul> <li>For diagnosis of Generalized Anxiety Disorder (GAD) or Social Anxiety Disorder (SAD): Require trial with a Selective-Serotonin Reuptake Inhibitor (SSRI) or a Serotonin-Norepinephrine Reuptake Inhibitor (SNRI) prior to initial benzodiazepine prescription</li> <li>For diagnosis of Panic Disorder: Require concurrent therapy with an antidepressant (SSRI, SNRI, or Tricyclic antidepressant [TCA]).</li> <li>For diagnosis of skeletal muscle spasms: Require trial with a skeletal muscle relaxant prior to a benzodiazepine</li> </ul>	DURATION LIMIT: > For Insomnia: 30 consecutive days > For Panic Disorder: 30 consecutive days	<ul> <li>Require confirmation of FDA approved or compendia supported use</li> <li>PA required for initiation of benzodiazepine therapy in patients currently on opioid or oral buprenorphine therapy</li> <li>PA required for any additional oral benzodiazepine prescription in patients currently on benzodiazepine therapy</li> </ul>
Crofelemer (Fulyzaq <sup>©</sup> )	Requires trial with an alternative anti- diarrheal agent		Confirm diagnosis of HIV/AIDS or antiretroviral therapy in daims history
Cyclosproine ophthalmic (Restasis <sup>6</sup> )	utilization as a first line agent or attempt	QUANTITY LIMIT: 60 vials dispensed as a 30-day supply	
Cystic fibrosis agents  ivacattor (Kalydeco)  ivacattor / lum acattor (Orkambi)			Confirmation of FDA-approved or compendia-supported indications     Genetic testing required to verify appropriate mutations
Dextromethorphan / quinidine (Nuedexta <sup>9</sup> )		QUANTITY LIMIT:  Two (2) capsules per day; 60 units per 30 days  DURATION LIMIT:  90 days of therapy	For patients ≥ 18 years of age: Requires confirm ation of diagnosis of P seudobulb ar affect

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Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Dronabinol (Marinol <sup>©</sup> )	<ul> <li>Step therapy for beneficiaries with HIV/ADS, or cancer, AND eating disorder: trial with megestrol acetate suspension prior to dronabinol</li> <li>Step therapy for beneficiaries with diagnosis of cancer and nausea /vomiting: trial with a NYS Medicaid-preferred 5-HT3 receptor antagonist prior to dronabinol</li> </ul>		Confirm diagnosis for Medicaid covered uses as follows:  > HIV/AIDS or Cancer and eating disorder  > Cancer and nausea/vomiting
Fentanyl transmucosal agents		QUANTITY LIMIT:  ➤ 4 units per day, 120 units per 30 days  DURATION LIMIT:  ➤ 90 days  ➤ Quantity and duration limits are not applicable to patients with a documented cancer or sickle cell diagnosis	<ul> <li>Limited to a total of four (4) opioid prescriptions every 30 days; Exemption for diagnosis of cancer or sickle cell disease</li> <li>For opioid-naïve patients - limited to a 15 days supply for all initial opioid prescriptions, Exemption for diagnosis of cancer or sickle cell disease</li> <li>Medical necessity rationale for opioid therapy is required for patients on established buprenorphine therapy</li> <li>PA is required for initiation of opioid therapy in patients currently on benzodiazepine therapy</li> </ul>
Irritable Bowel Agents  ➤ linaclotide (Linzess™)  ➤ lubiprostone (Amitiza®)*  *(lubiprostone parameters for opioid induced constipation see Opioid induced constipation agents section below)	Step therapy with trials of both a bulking- agent and an osmotic laxative prior (defined as within 89 days) to lubiprostone or linadotide	DURATION LIMIT: ➤ 30 days with 2 refills/prescription	
Lipid Lowering Agents - Proprotein Convertase Subtilisin Kexin 9 (PCSK9) Inhibitors ➤ alirocumab (Praluent™) ➤ evolocumab (Repatha™)	Require trial of a HMG-CoA Reductase Inhibitors (Statin) at maximum tolerated dosage		Confirm diagnosis for the FDA-approved indication of.  Familial hypercholesterolemia (heterozygous or homozygous)  Atherosclerotic cardiovascular disease Require concurrent statin therapy
Lipid Lowering Agents - Triglyceride transfer protein inhibitors:  Iomitapide (Juxtapid <sup>®</sup> ) mipomersen (Kynamro <sup>®</sup> )	Requires trial with high intensity statin therapy		Confirm diagnosis of homozygous familial hypercholesterolemia
Metozolv <sup>®</sup> ODT (metodopramide)	Requires a trial with conventional metoclopramide before metoclopramide orally disintegrating tablet (ODT), except with diagnosis of diabetes mellitus	QUANTITY LIMIT: ➤ 4 units per day, 120 units per 30 days  DURATION LIMIT: ➤ 90 days	

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Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Methadone		QUANTITY LIMIT:  > 12 units per day, 360 units per 30 days  > Exemption for diagnosis of cancer or sickle cell disease	➤ Limited to a total of four (4) opioid prescriptions every 30 days; Exemption for diagnosis of cancer or sickle cell disease     ➤ Medical necessity rationale for methadone is required for patients on established buprenorphine therapy     ➤ PA required for methadone prescriptions for patients currently on long-acting opioid therapy. Exemption for diagnosis of cancer or sickle cell disease     ➤ PA required for initiation of long-acting opioid therapy in opioid-naïve patients. Exemption for diagnosis of cancer or sickle cell disease     ➤ PA required for initiation of methadone therapy in patients currently on benzodiazepine therapy
Metreleptin (Myalept <sup>®</sup> )			Confirm diagnosis for the FDA-approved indications:  Leptin deficiency in patients with congenital generalized lipodystrophy (CGL) OR  acquired generalized lipodystrophy (AGL)
Olanzapine / Fluoxetine (Symbyax <sup>®</sup> )	When prescribing for the treatment of major depressive disorder (MDD) in the absence of other psychiatric comorbidities, trial with at least one different antidepressant agent is required		PA is required for the initial prescription for beneficiaries younger than 18 years
Opioid Induced Constipation Agents  > lubiprostone (Amitiza®)*  > methylnaltrexone (Relistor®)  > naloxegol (Movantik™)  *(for lubiprostone parameters for irritable bowel, see Irritable bowel agents section above)	Trial with an osmotic laxative, a stim ulant laxative and a stool softener prior to use of lubiprostone, methylnaltrexone, or naloxegol	QUANTITY LIMIT:  ➤ lubiprostone: 2 capsules per day, 60 capsules per 30 days  ➤ methylnaltrexone: 1 vial or syringe per day (30 vials/syringes per 30 days, 4 kits per 28 days  ➤ naloxegol: 1 tablet per day, 30 tablets per 30 days	Confirmation of FD A-approved or compendia- supported indications
Oral Pollen/Allergen Extracts (Grastek <sup>6</sup> , Oralair <sup>6</sup> , Ragwitek <sup>6</sup> )	Trial with a preferred intranasal corticosteroid		Confirm diagnosis for the FDA-approved indication of Pollen-induced allergic rhinitis confirmed by positive skin or in vitro testing for pollen-specific IgE antibodies
Pyrimethamine (Daraprim <sup>®</sup> )			Confirmation of FDA-approved or compendia-supported indications     Require concurrent utilization of leucovorin

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Drug / Class Name	Step Therapy (ST) Parameters	Frequency / Quantity / Duration (F/Q/D) Parameters	Additional / Alternate Parameter(s)
Pulmonary Fibrosis Agents  > Ofev <sup>6</sup> > Esbriet <sup>6</sup>			Confirm diagnosis for the FDA-approved indication of treatment of idiopathic pulmonary fibrosis (IPF)
Quinine		QUANTITY AND DURATION LIMITS:  > Maximum 42 capsules as a 7-day supply > limited to 1 prescription per year	
Tasimetteon (Hetlioz <sup>®</sup> )		QUANTITY LIMIT: > One unit per day, 30 units per 30 days	Confirm diagnosis of Non-24-hour sleep-wake disorder in totally blind patients
Ta zarotene (Tazorac <sup>6</sup> )			Confirm diagnosis for Medicaid covered uses
Tetrabenazine (Xenazine <sup>®</sup> )			Confirm diagnosis of one of the following FDA and Compendia approved indication in patients ≥ 18 years of age:
			<ul> <li>Chorea associated with Huntington's disease</li> </ul>
			➤ Gilles de la Tourette's syndrome ➤ Tardive dyskinesia
Teriparatide (Forteo <sup>9</sup> )	Requires a trial with a preferred oral bisphosphonate prior to teriparatide	QUANTITY LIMIT:  > One unit (2.4 mL) per 30-day period	
		LIFE TIME QUANTITY LIMIT:	
		➤ 25 months of therapy	

For more information on DUR Program, please refer to <a href="http://nyhealth.gov/health\_care/medicaid/program/dur/index.htm">http://nyhealth.gov/health\_care/medicaid/program/dur/index.htm</a>.

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## NYS Medicaid Fee-For-Service Brand Less Than Generic (BLTG) Program

On April 26, 2010, New York Medicaid implemented a new cost containment initiative, which promotes the use of certain multi-source brand name drugs when the cost of the brand name drug is less expensive than the generic equivalent.

In conformance with State Education Law, which intends that patients receive the lower cost alternative, brand name drugs included in this program:

- Do not require "Dispense as Written" (DAW) or "Brand Medically Necessary" on the prescription
- Have a generic copaγment
- Are paid at the Brand Name Drug reimbursement rate or usual and customary price, whichever is lower
- . Do not require a new prescription if the drug is removed from this program

### Effective February 18, 2016:

- Gleevec will be added to the program.
- Bactroban cream, Kadian, and Metrogel will be removed from the program.

### Current list of Brand name drugs included in this program\* (Updated 2/03/2016):

\*List is subject to change

Abilify tablet	Combivir	Mepron	Tobradex suspension
Adderall XR	Copaxone 20 mg/mL SubQ	Myfortic	Tricor
Aggrenox	Aggrenox Diastat Niaspan		Trilipix
Aldara	Epivir HBV tablet	Patanase	Trizivir
Alphagan P0.15%	Exelon Patch	Protopic	Valcyte
Astepro	Astepro Focalin XR 5mg, 10 mg, 15mg, 20mg, 30mg, 40mg		Wellbutrin
Baraclude	Gabitril 2mg, 4mg	Soriatane	Xeloda
Catapres-TTS	Gleevec	Tegretol suspension	Xenazine
Cellcept suspension	Hepsera	TegretoIXR	

Please keep in mind that drugs in this program may be subject to prior authorization requirements of other pharmacy programs; again promoting the use of the most cost-effective product.

### IMPORTANT BILLING INFORMATION

- Prescription claims submitted to the Medicaid program DO NOT require the submission of Dispense As Written/Product Selection Code of '1';
- · Pharmacies can submit any valid NCPDP field (408-D8) value

For more information on the Brand Less Than Generic (BLTG) Program, please refer to https://newyork.fhsc.com/providers/bltqp\_about.asp.

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## NYS Medicaid Fee-For-Service Mandatory Generic Drug Program

State law excludes Medicaid coverage of brand name drugs that have a Federal Food and Drug Administration (FDA) approved A-rated generic equivalent, unless a prior authorization is obtained.

Coverage parameters under the Preferred Drug Program (PDP), Clinical Drug Review Program (CDRP), and/or the Brand Less Than Generic (BLTG) Program are applicable for certain products subject to the Mandatory Generic Drug Program (MGDP), including exemptions (as listed below).

### **Prior Authorization Process**

- Prescribers, or an agent of the prescriber, must call the prior authorization line at 1-877-309-9493 and respond to a series of questions that
  identify the prescriber, the patient and the reason for prescribing this drug. The Mandatory Generic Program Prescriber Worksheet and
  Instructions provide step-by-step assistance in completing the prior authorization process.
- The prescriber must write "DAW and Brand Medically Necessary" on the face of the prescription.
- The call line 1-877-309-9493 is in operation 24 hours a day, seven days a week.

### Exempt Drugs

 Based on specific characteristics of the drug and/or disease state generally treated, the following brand name drugs are exempt from the program and do NOT require PA:

Clozaril <sup>®</sup>	Levothyroxine Sodium (Unithroid <sup>®</sup> , Synthroid <sup>®</sup> , Levoxyl <sup>®</sup> )
Coumadin <sup>®</sup>	Neoral <sup>®</sup>
Dilantin <sup>®</sup>	Sandimmune <sup>®</sup>
Gengraf <sup>®</sup>	T egretol®
Lanoxin <sup>®</sup>	Zarontin <sup>®</sup>

For more information on the Mandatory Generic Program, please refer to https://newyork.fhsc.com/providers/MGDP\_about.asp.

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## NYS Medicaid Fee-For-Service Dose Optimization Program

On November 14, 2013, the Medicaid Fee-for-Service program instituted a Dose Optimization initiative. Dose optimization can reduce prescription costs by reducing the number of pills a patient needs to take each day. The Department has identified drugs to be included in this program, the majority of which have FDA approval for once-a-day dosing, have multiple strengths available in correlating increments at similar costs and are currently being utilized above the recommended dosing frequency. Prior authorization will be required to obtain the following medication beyond the following limits:

### **Dose Optimization Chart**

Brand Name Dose Optimization Limitations							
CARDIOVASCULAR							
Angiotensin Receptor Blockers (ARBs)							
Benicar 20m g	1 daily	Tablet					
Micardis 20mg, 40mg	1 daily	Tablet					
Diovan 40mg, 80mg, 160mg	1 daily	Tablet					
	ARBs/ (	Calcium Channel B	llockers				
Exforge 5–160mg	1 daily	Tablet					
		ARBs/ Diuretics					
Benicar HCT 20-12.5mg	1 daily	Tablet					
Diovan HCT 80-12.5mg, 160-12.5mg	1 daily	Tablet					
Edarbydor 40–12.5mg	1 daily	Tablet					
Micardis HCT 40-12.5mg, 80-12.5mg	1 daily	Tablet					
		Beta Blockers					
Bystolic 2.5mg, 5mg, 10mg	1 daily	Tablet					
Coreg CR 20mg,40mg	1 daily	Tablet					
nadolol 40mg	1 daily	Tablet					
Toprol XL 25mg, 50mg, 100mg	1 daily	Tablet					
	HMG C	o A Reductase Inf	ibitors				
Crestor 5mg, 10mg, 20mg	1 daily	Tablet					
		Niacin Derivatives					
Niaspan 500mg	1 daily	Tablet					
	Anticonvu	ılsants – Second (	eneration				
Lyrica 25mg, 50mg, 75mg, 100mg, 150mg, 200mg	3 daily	Capsule	Electronic bypass for diagnosis of seizure disorder indentified in medical				
Lyrica 225mg and 300mg	2 daily	Capsule	daims data.				

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Brand Name Dose Optimization Limitations							
Diam name	CENTRAL NERVOUS SYSTEM						
Antiparkinson Agents							
Azilect 0.5mg 1 daily Tablet							
Antipsychotics – Second Generation							
Ability 2mg	4 daily	Tablet					
Abilify 5mg, 10mg, 15mg	1 daily	Tablet	In the case of dose titration for these once daily medications, the Department				
Invega 1.5mg, 3mg	1 daily	Tablet	will allow for multi-day dosing (up to 2 doses/daily) for titration purposes for 3				
Latuda 20mg, 40mg, 60mg	1 daily	Tablet	months.				
olan zapine 5mg	1 daily	Tablet	1				
olan zapine ODT 5m g	1 daily	Tablet					
Seroquel XR 150mg, 200mg	1 daily	Tablet					
Sym byax 3–25m g, 6–25m g, 12–25m g	1 daily	Capsule					
Zyprexa Zydis 5mg, 10mg	1 daily	Tablet					
		CNS Stimulants					
Concerta ER 18mg, 27mg, 54mg	1 daily	Tablet					
Concerta ER 36mg	2 daily	Tablet					
Focalin XR 5mg, 10mg, 15mg, 20mg	1 daily	Capsule					
Metadate CD 10mg, 20mg	1 daily	Capsule					
Provigil 100mg	1 daily	Tablet					
QuillichewER 20mg, 40mg	1 daily	Tablet					
QuillichewER 30mg	2 daily	Tablet					
Ritalin LA 10mg, 20mg	1 daily	Capsule					
Vyvanse 20mg, 30mg	1 daily	Capsule					
	Non-Ergot	Dopamine Recept	or Agonists				
Requip XL 2mg, 4mg, 6mg	1 daily	Tablet					
	Other Agents for Atten	tion Deficit Hypera	ctivity Disorder (ADHD)				
guanfacine ER 1mg, 2mg, 3 mg, 4mg	1 daily	Tablet					
Intuniv1mg,2mg	1 daily	Tablet					
Strattera 40mg	1 daily	Capsule					
	<u> </u>	Sedative Hypnotics	s				
Lunesta 1mg	1 daily	Tablet					
	•	•					

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Brand Name	Dose Optimization Limitations						
CENTRAL NERVOUS SYSTEM							
Serotonin-Norepinephrine Reuptake Inhibitors (SNRIs)							
Effexor XR 37.5mg, 75mg	1 daily	Capsule	In the case of dose titration for these once daily medications, the Department				
Pristiq ER 50mg	1 daily	Tablet	will allow for multi-day dosing (up to 2 doses/daily) for titration purposes for three months.				
Selective Serotonin Reuptake Inhibitors (SSRIs)							
Lexapro 5mg, 10mg	1 daily	Tablet	In the case of dose titration for these once daily medications, the Department				
Viibryd 1 0mg, 20mg	1 daily	Tablet	will allow for multi-day dosing (up to 2 doses/daily) for titration purposes for three months.				
	ENDO	CRINE AND META	BOLIC				
	Dipeptidyl I	Peptidase-4 (DPP-4	1) Inhibitors				
Januvia 25mg, 50mg	1 daily	Tablet					
Onglyza 2.5m g	1 daily	Tablet					
	Thia	azolidinediones (T	ZDs)				
Actos 15mg	1 daily	Tablet					
Actoplus Met XR 15-1000mg	1 daily	Tablet					
	G	ASTROINTESTINA	IL .				
	Pr	oton Pump Inhibite	prs				
Dexilant 30mg	1 daily	Capsule					
Nexium 20mg	1 daily	Capsule					
Prevacid DR 15mg	1 daily	Capsule					
	RENA	L AND GENITOUR	INARY				
	Urina	ry Tract Antispasn	nodics				
Detrol LA 2mg	1 daily	Capsule					
Enablex 7.5mg	1 daily	Tablet					
oxybutynin chloride ER 5mg	1 daily	Tablet					
Toviaz ER 4mg	1 daily	Tablet					
Vesicare 5mg	1 daily	Tablet					

PA requirements are not dependent on the date a prescription is written. New prescriptions and refills on existing prescriptions require PA even if the prescription was written before the date the drug was determined to require PA.

To obtain a prior authorization (PA), please call the prior authorization Clinical Call Center at 1-877-309-9493. The Clinical Call Center is available 24 hours per day, 7 days per week with pharmacy technicians and pharmacists who will work with you, or your agent, to quickly obtain PA.

Medicaid enrolled prescribers with an active e-PACES account can initiate PA requests through the web-based application PAX press<sup>®</sup>. The website for PAX press is <a href="https://paxpress.nypa.hidinc.com">https://paxpress.nypa.hidinc.com</a>.

## 5 - Preferred Supply List (as of March 2016)

### **NYS Diabetic Supplies**

	WTS DIabetic Supp	mes	n I
he co			Revised: 10/01/2015
Manufacturer	Product	NDC	STRIPS/ METERS
Abbott	FreeStyle Lite Meter	99073070805	Meter
Abbott	FreeStyle Lite Test Strips - 50ct	99073070822	Strips
Abbott	FreeStyle Lite Test Strips - 100ct	99073070827	Strips
Abbott	FreeStyle Freedom Lite Meter	99073070914	Meter
Abbott	FreeStyle InsuLinx Test Strips - 50ct	99073071231	Strips
Abbott	FreeStyle In suLinx Meter	99073071143	Meter
Abbott	FreeStyle InsuLinx Test Strips - 100 ct	99073071227	Strips
Abbott	Precision Xtra Beta Ketone Test Strips- 10ct	57599074501	Strips
Bayer	BREEZE Blood Glucose Meter	00193144001	Meter
Bayer	BREEZE 2 Test Strip - 50ct	00193146550	Strips
Bayer	BREEZE 2 Test Strip - 100 ct	00193146621	Strips
Bayer	CONTOUR Test Strips - 50ct	00193708050	Strips
Bayer	CONTOUR Test Strips - 100 ct	00193709021	Strips
Bayer	CONTOUR Blood Glucose Meter	00193715101	Meter
Bayer	CONTOUR NEXT EZ Blood Glucose Meter	00193725201	Meter
Bayer	CONTOUR NEXT Test Strips - 50ct	00193731150	Strips
Bayer	CONTOUR NEXT Test Strips - 100ct	00193731221	Strips
Bayer	CONTOUR NEXT Blood Glucose Meter	00193737701	Meter
Bayer	CONTOUR NEXT USB Mieter	00193739301	Meter
Bayer	CONTOUR NEXT USB Blood Glucose Meter	00193741101	Meter
LifeScan	One Touch UltraMini Meter - Silver Moon	53885020801	Meter
LifeScan	One Touch Ultra Blue Test Strips - 50ct	53885024450	Strips
LifeScan	One Touch Ultra Blue Test Strips - 100 ct	53885024510	Strips
LifeScan	One Touch Verio Test Strips - 25ct	53885027025	Strips
LifeScan	One Touch Verio Test Strips - 50ct	53885027150	Strips
LifeScan	One Touch Verio Test Strips - 100ct	53885027210	Strips
LifeScan	One Touch UltraMini Meter - Pink Glow	53885041901	Meter
LifeScan	One Touch UltraMini Meter - Limelight	53885042001	Meter
LifeScan	One Touch Ultra 2 Meter	53885044801	Meter
LifeScan	One Touch Verio Meter System	53885065701	Meter
LifeScan	One Touch UltraMini Meter - Blue Comet	53885091101	Meter
LifeScan	One Touch UltraMini Meter - Purple Twlight	53885091201	Meter
LifeScan	On e Touch Ultra Blue Test Strips - 25ct	53885099425	Strips
LifeScan	One Touch Verio IQ Meter	53885026701	Meter
Miedisense (Abbott)	Precision Xtra Meter	57599881401	Meter
Miedisense (Abbott)	Precision Xtra Test Strips - 50ct	57599972804	Strips
Miedisense (Abbott)	Precision Xtra Test Strips - 100 ct	5 <i>7</i> 599987705	Strips
Therasense(Abbott)	FreeStyle Test Strips - 50ct	99073012050	Strips
Therasense(Abbott)	FreeStyle Test Strips - 100 ct	99073012101	Strips

## 6 - Enrollee Brochure (English Version)

### **PDP**

### **New York State** Medicaid Preferred Drug Program

A GUIDE FOR PEOPLE WITH MEDICAID



What is the Medicaid Preferred Drug Program (PDP)?

This program encourages doctors to prescribe certain drugs, Called "preferred" drugs. When they prescribe other similar drugs which are not included on the preferred drug list, they need to get special approval (prior authorization) before you can receive the drug.

#### Who decides which drugs are "preferred"?

A committee made up of doctors, pharmacists, and patient advocates works with the Department of Health to review drugs and identify those that are safe, effective and less expensive. Preferred drugs have been found to be as effective as non-preferred drugs.

#### What if I don't want to change my medications?

Only your doctor can decide which drugs you should take. Ask your doctor or pharmacist if you have questions about changes made to your prescriptions.

Need help? Call the Medicaid Help 1-800-541-2831



### Remember:

- All drugs that Medicaid currently covers are still available.
- · Only your doctor can decide which drugs you should take.
- Ask your doctor or pharmacist if you have questions about your medicine.

### What if I need my medication and the doctor's office is closed?

If your doctor cannot be contacted, and you have a valid prescription, the pharmacist can give you a 72-hour emergency supply of medicine until your doctor can be contacted.

For more information, visit the NYS Medicaid Preferred Drug Program Website: https://newyork.fhsc.com



### **MGDP**

### **New York State** Medicaid Generic Drug Program

A GUIDE FOR PEOPLE WITH MEDICAID AND FAMILY HEALTH PLUS



What is the Generic Drug Program? The law requires doctors to prescribe the generic version of a drug, unless they get special approval for a brand name drug.

### What is a generic drug?

A generic drug is a copy of a brand name drug.

It is the same medicine with the same active ingredients as the brand name drug, but usually

## Is a generic drug as good as a brand name drug?

Yes. The federal government makes certain that the generic drug is as safe and effective as the brand name drug. (You may already be taking generic drugs).

What if I am taking a brand name drug that has a generic version? Medicaid will not pay for your brand name drug unless your doctor calls Medicaid to get approval, and writes the approval number

1-800-541-2831



### Remember:

- Only your doctor can decide which drugs you should take.
- Generic drugs are safe and effective copies of brand name drugs and are approved by the federal government.
- Ask your doctor and pharmacist about generic drugs.

What if my doctor forgets to get the approval for my brand name drug? The pharmacist can call your doctor to discuss if the generic drug is right for you

What if I really need my medicine and the doctor's office is closed? In an emergency, if you have a valid prescription, the pharmacist may give you a small supply of the brand name drug until you can talk to someone at your doctor's office or clinic.

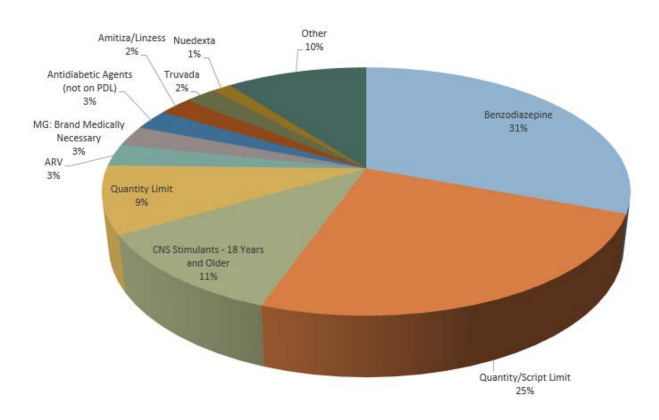
Why are my pills a different color than they used to be? Generic pills may look different because they are made by another company. They may be a different color or shape, but they are as safe and effective as the brand name drug.



## 7 – Preferred Drug Program Website Information

- Information about the NY Medicaid Pharmacy Prior Authorization Programs can be accessed on the Internet at: <a href="https://newyork.fhsc.com/">https://newyork.fhsc.com/</a> or <a href="https://www.health.state.ny.us">https://www.health.state.ny.us</a>
- The complete PDL can be accessed at: <a href="https://newyork.fhsc.com/downloads/providers/NYRx">https://newyork.fhsc.com/downloads/providers/NYRx</a> PDP PDL.pdf

## 8 - CDRP and Other Prior Authorizations by Type

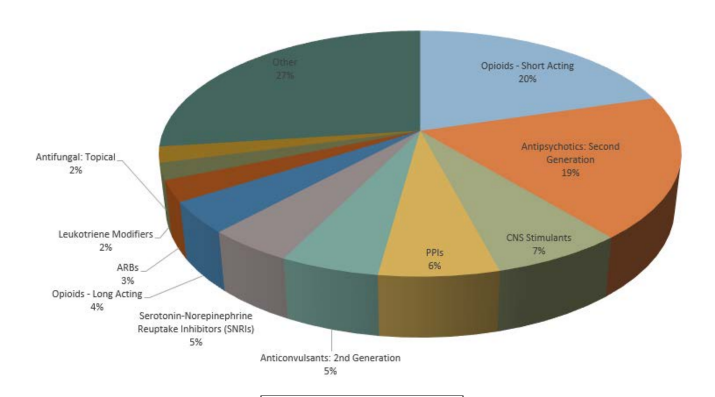


\*\*This chart represents Approved PAs for the following: drugs/drug classes subject to step therapy, FQD (Frequency, Quantity and Duration Limits), PDP classes subject to CDRP and CDRP.

Total PAs = 50,940

1	Benzodiazepine	15665	25	Xenazine	52
2	Quantity/Script Limit	12560	26	Actiq/Fentora	52
3	CNS Stimulants - 18 Years and Older	5680	27	Dose Optimiz	49
4	Quantity Limit	4508	28	Forteo	46
5	ARV	1446	29	Progesterone	33
6	MG: Brand Medically Necessary	1375	30	MG: Generic	32
7	Antidiabetic Agents (not on PDL)	1336	31	CF Agents	25
8	Amitiza/Linzess	1245	32	Movantik	24
9	Truvada	1155	33	Acthar	22
10	Nuedexta	714	34	Regranex	14
11	Lidoderm	636	35	Growth Horm	14
12	BLTG	597	36	Xyrem	12
13	Anabolic Steroids	596	37	Opioid/Bupre	11
14	Synagis	526	38	Serostim	4
15	Restasis	389	39	PCSK9 Inhibi	4
16	Methadone	354	40	Pulmonary Fi	4
17	Marinol	332	41	Quinine	3
18	DUR: Drug to Drug Interaction	292	42	Metozolv	2
19	Immunomodulators: Topical	282	43	Fulyzaq	2
20	Antifungals: Topical Onychomycosis	252	44	Oral Pollen/A	2
21	Oxazolidinone Antibiotics	235	45	Daraprim	2
22	Inhaled Antibiotics for CF	177	46	Hetlioz	2
23	PDE-5 Inhibitorsfor Pulmunoary Hypertension	118	47	Script Limit	1
24	Tazorac	57	48	Biotin	1

## 9 – PDP Prior Authorizations by Class



Total PDP PAs = 106,319

Of the PAs issued in this fiscal year, the following PDP drug classes are listed by the number of PAs requested:

Opioids - Short Acting	21635	Antibiotics: GI	334	Inh. Long Acting Beta-2 Adrenergic	64
Antipsychotics: Second Generation	19838	Multiple Sclerosis Agents	312	Antifungals: Oral	56
CNS Stimulants	7341	Triptans	304	Antivirals: Oral	55
PPIs	6633	Ophthalmics: Prostaglandin Agonists	295	Antibiotics: Topical	52
Anticonvulsants: 2nd Generation	5477	Topical Steroids: Low Potency	293	Non-Ergot Dopamine Receptor Agonis	t 49
Serotonin-Norepinephrine Reuptake Inhibitors (SNRIs)	4811	Cholesterol Absorption Inhibitors	279	Alpha-Glucosidase Inhibitors	44
Opioids - Long Acting	4348	Ophthalmics: Antihistamines	264	Platelet Inhibitors	43
ARBs	2927	Carbamazepine Derivatives	255	Inhaled Corticosteroids	29
Leukotriene Modifiers	2305	Topical Steroids: Medium Potency	237	Ophthalmics: NSAIDs	29
Antifungal: Topical	2229	Antihistamines: Nasal	224	Otics: Quinolones	29
NSAIDs: Rx	2059	Fluoroquinolones	208	Ophthalmic Antibiotic/Steroid Combo	28
Other Agents for ADHD	1616	Anticholinergics/COPD Agents	205	Calcium Channel Blockers (DHP)	25
DPP-4 Inhibitors	1489	Alzheimer's Agents	200	Antiemetics	23
Hep C: Direct Acting Antivirals	1458	Tetracycline	185	Pancreatic Enzymes	23
Sedative Hypnotics	1357	Thiazolidinediones	185	Psoriasis Agents: Topical	23
Urinary Tract Antispasmodics	1256	Bisphosphonates	167	Progestins	20
Inh. Short Acting Beta-2 Adrenergic	1245	Biguanides	166	Actinic Keratosis Agents	19
GLP-1 Agonist	949	Antivirals: Topical	148	Benzodiazepines: Rectal	16
Triglyceride Agents	937	Growth Hormones	140	PAH Oral Agents - Other	14
Antihistamines: 2nd Generation	909	GI Prep Agents	139	Ophthalmics: Alpha-2 Adrenergics	14
Statins	827	Ophthalmics: Quinolones	136	Beta Blocker/Diuretic Combinations	11
Beta Blockers	799	Topical Steroids: Very High Potency	128	Hepatitis B Agents	11
ARB Combinations	753	Glucocorticoid: Oral	127	Niacin Derivatives	9
Anticoagulants: Oral	724	Erythropoiesis Stimulating Agents (ESAs)	117	Insulin: Rapid Acting	8
Skeletal Muscle Relaxants	697	Xanthine Oxidase Inhibitors	113	Antipsychotics: Injectable	7
Selective Serotonin Reuptake Inhibitors (SSRIs)	658	ACE Inhibitors	100	Direct Renin Inhibitors	6
Immunomodulators: Systemic	610	Meglitinides	100	Hepatitis C Agents: Injectable	6
Opioid Dependence Agents	601	Ophthalmics: Antibiotics	100	H. Pylori Agents	5
Antiinfectives: Topical	513	Inhaled Antibiotics	97	Opioid Antagonists	5
Sulfasalazine Derivatives	468	ARB/CCB Combinations	95	ACE Inhibitor Combinations	4
Topical Steroids: High Potency	455	Alpha Reductase Inhibitor: BPH	88	ACE Inhibitor/Diuretic Combinations	3
ARB/Diuretic Combinations	446	Inhaled Steroid/Beta2 LA Combo	86	Amylin Analog	3
SGLT2 Inhibitors	404	Anticoagulants: Injectable	79	Cephalosporins: Third Generation	3
Phosphate Binders/Regulators	391	Insulin: Long Acting	68	Cystine Depleting Agents	2
Steroids: Intranasal	373	Selective Alpha Adrenergic Blockers	68	Ophthalmics: Beta Blockers	1

# 10 – PDP and Diabetic Supply Cost Avoidance by County

County	PDP	Diabetic Supplies	Total	% Total
Albany	-\$414,437	\$57,421	-\$357,017	1.12%
Allegany	-\$116,285	\$26,969	-\$89,316	0.28%
Broome	-\$332,631	\$43,588	-\$289,043	0.91%
Cattaraugus	-\$170,680	\$22,491	-\$148,189	0.46%
Cayuga	-\$139,495	\$17,117	-\$122,379	0.38%
Chautauqua	-\$218,854	\$28,362	-\$190,492	0.60%
Chemung	-\$210,288	\$32,641	-\$177,647	0.56%
Chenango	-\$117,295	\$23,287	-\$94,008	0.29%
Clinton	-\$220,986	\$40,204	-\$180,781	0.57%
Columbia	-\$103,915	\$9,952	-\$93,964	0.29%
Cortland	-\$73,954	\$4,976	-\$68,978	0.22%
Delaware	-\$172,903	\$32,641	-\$140,262	0.44%
Dutchess	-\$362,021	\$37,119	-\$324,901	1.02%
Erie	-\$1,153,092	\$265,608	-\$887,484	2.78%
Essex	-\$90,638	\$11,245	-\$79,392	0.25%
Franklin	-\$198,602	\$35,030	-\$163,572	0.51%
Fulton	-\$128,115	\$12,440	-\$115,675	0.36%
Genesee	-\$90,556	\$9,155	-\$81,401	0.25%
Greene	-\$61,350	\$6,867	-\$54,483	0.17%
Hamilton	-\$6,945	\$796	-\$6,149	0.02%
Herkimer	-\$108,973	\$30,949	-\$78,024	0.24%
Jefferson	-\$301,834	\$32,144	-\$269,690	0.84%
Lewis	-\$44,513	\$4,180	-\$40,334	0.13%
Livingston	-\$85,957	\$9,653	-\$76,304	0.24%
Madison	-\$113,929	\$8,359	-\$105,570	0.33%
Monroe	-\$1,278,127	\$288,596	-\$989,531	3.10%
Montgomery	-\$90,760	\$11,842	-\$78,918	0.25%
Nassau	-\$1,100,074	\$164,500	-\$935,574	2.93%
Niagara	-\$276,614	\$36,025	-\$240,590	0.75%
Oneida	-\$433,640	\$66,576	-\$367,063	1.15%
Onondaga	-\$828,875	\$111,259	-\$717,616	2.25%
Ontario	-\$119,487	\$9,952	-\$109,536	0.34%
Orange	-\$486,056	\$42,792	-\$443,264	1.39%
Orleans	-\$70,507	\$6,170	-\$64,337	0.20%
Oswego	-\$156,923	\$23,287	-\$133,637	0.42%
Otsego	-\$116,357	\$8,359	-\$107,997	0.34%
Putnam	-\$57,832	\$1,990	-\$55,841	0.17%
Rensselaer	-\$218,140	\$26,471	-\$191,669	0.60%
Rockland	-\$437,403	\$43,489	-\$393,914	1.23%
St. Lawrence	-\$442,808	\$56,824	-\$385,984	1.21%
Saratoga	-\$212,929	\$23,685	-\$189,245	0.59%
Schenectady	-\$226,085	\$39,309	-\$186,776	0.58%
Schoharie	-\$42,647	\$7,464	-\$35,183	0.11%
Schuyler	-\$38,986	\$2,985	-\$36,001	0.11%

County	PDP	Diabetic Supplies	Total	% Total
Seneca	-\$50,856	\$7,563	-\$43,293	0.14%
Steuben	-\$301,018	\$40,204	-\$260,813	0.82%
Suffolk	-\$1,489,354	\$146,189	-\$1,343,164	4.21%
Sullivan	-\$161,767	\$15,624	-\$146,143	0.46%
Tioga	-\$112,247	\$16,520	-\$95,727	0.30%
Tompkins	-\$132,489	\$14,131	-\$118,358	0.37%
Ulster	-\$233,865	\$25,974	-\$207,892	0.65%
Warren	-\$152,865	\$14,927	-\$137,937	0.43%
Washington	-\$103,630	\$11,345	-\$92,285	0.29%
Wayne	-\$128,900	\$25,476	-\$103,424	0.32%
Westchester	-\$1,022,673	\$222,816	-\$799,857	2.51%
Wyoming	-\$100,693	\$19,406	-\$81,287	0.25%
Yates	-\$28,982	\$3,384	-\$25,599	0.08%
Sub Totals	-\$15,691,837	\$2,338,328	-\$13,353,509	41.82%
New York City	-\$21,021,518	\$3,478,185	-\$17,543,333	54.95%
ОМН	-\$439,809	\$51,848	-\$387,962	1.22%
OMR	-\$559,552	\$46,972	-\$512,580	1.61%
NYS DOH	-\$156,862	\$26,670	-\$130,192	0.41%
Grand Total	-\$37,869,578	\$5,942,003	-\$31,927,575	