

### BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA

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EFFECTIVE 04/01/12 Version 2012.5a

- Prior authorization for a non-preferred agent in any category will be given only if there has been a trial of the preferred brand/generic equivalent or preferred formulation of the active ingredient, at a therapeutic dose, that resulted in a partial response with a documented intolerance.
- Prior authorization of a non-preferred isomer, pro-drug, or metabolite will be considered with a trial of a preferred parent drug of the same chemical entity, at a therapeutic dose, that resulted in a partial response with documented intolerance or a previous trial and therapy failure, at a therapeutic dose, with a preferred drug of a different chemical entity indicated to treat the submitted diagnosis. (The required trial may be overridden when documented evidence is provided that the use of these preferred agent(s) would be medically contraindicated.)
- Unless otherwise specified, the listing of a particular brand or generic name includes all legend forms of that drug. OTC drugs are not covered unless specified.
- PA criteria for non-preferred agents apply in addition to general Drug Utilization Review policy that is in effect for the entire pharmacy program, including, but not limited to, appropriate dosing, duplication of therapy, etc.
- The use of pharmaceutical samples will not be considered when evaluating the members' medical condition or prior prescription history for drugs that require prior authorization.
- Acronyms
  - CL Requires clinical PA. For detailed clinical criteria, please refer to: <u>http://www.dhhr.wv.gov/bms/Pharmacy/Pages/PriorAuthorizationCriteria.aspx</u>
  - NR New drug has not been reviewed by P & T Committee
  - AP Non-preferred and selected preferred drugs, where indicated, are subject to auto-PA criteria. See PA criteria column.



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|-----------------|---|---|--|--|--|
| DRUG CLASS      |   |   |  |  |  |
| ACNE AGENTS (To | ACNE AGENTS (Topical) <sup>AP</sup>   |   |  |  |  |
|                 |   | FECTIVE   |  |  |  |
|                 | AKNE-MYCIN (erythromycin)<br>AZELEX (azelaic acid)<br>clindamycin<br>erythromycin<br>sodium sulfacetamide | ACZONE (dapsone)<br>CLEOCIN-T (clindamycin)<br>EVOCLIN (clindamycin)<br>KLARON (sodium sulfacetamide) | Thirty (30) day trials each of one<br>preferred retinoid and two unique<br>chemical entities in two other<br>subclasses, including the generic<br>version of a requested non-<br>preferred product, are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present. (In cases of pregnancy, a<br>trial of retinoids will not be<br>required.) |  |  |
|                 | RETIN   | NOIDS   |  |  |  |
|                 | RETIN A liquid & Micro (tretinoin)  | adapalene   | PA required after 17 years of age  |  |  |
|                 | TAZORAC (tazarotene)  | AVITA (tretinoin)   | for tretinoin products.  |  |  |
|                 | tretinoin cream, gel  | DIFFERIN (adapalene)  |  |  |  |
|                 |   | RETIN-A cream, gel (tretinoin)  |  |  |  |
|                 |   | TRETIN-X (tretinoin)  |  |  |  |
|                 | KERATOLYTICS (  | Senzoyl Peroxides)  |  |  |  |
|                 | benzoyl peroxide  | BENZAC WASH (benzoyl peroxide)  | Acne kits are non-preferred.   |  |  |
|                 | ETHEXDERM (benzoyl peroxide)  | BENZEFOAM (benzoyl peroxide)  |  |  |  |
|                 | OSCION (benzoyl peroxide)   | BENZEFOAM ULTRA (benzoyl peroxide)  |  |  |  |
|                 |   | BREVOXYL (benzoyl peroxide)   |  |  |  |
|                 |   | DESQUAM (benzoyl peroxide)  |  |  |  |
|                 |   | LAVOCLEN (benzoyl peroxide)   |  |  |  |
|                 |   | TRIAZ (benzoyl peroxide)  |  |  |  |
|                 | COMBINATION AGENTS  |   |  |  |  |
|                 | benzoyl peroxide/urea   | ACANYA (clindamycin phosphate/benzoyl   | Thirty (30) day trials each of one   |  |  |
|                 | erythromycin/benzoyl peroxide   | peroxide)   | preferred retinoid and two unique  |  |  |
|                 | sulfacetamide sodium/sulfur wash/cleanser   | AVAR (sulfur/sulfacetamide)   | chemical entities in two other   |  |  |
|                 |   | BENZACLIN GEL (benzoyl peroxide/  | subclasses, including the generic  |  |  |
|                 |   | clindamycin)  | version of a requested non-  |  |  |
|                 |   | BENZAMYCIN PAK (benzoyl peroxide/   | preferred product, are required<br>before a non-preferred agent will be  |  |  |
|                 |   | erythromycin)   | authorized unless one of the   |  |  |
|                 |   | benzoyl peroxide/clindamycin gel  | exceptions on the PA form is   |  |  |
| 1               | I   |   | -  |  |  |



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|                           |                  | CLENIA (sulfacetamide sodium/sulfur)<br>DUAC CS (benzoyl peroxide/ clindamycin)<br>EPIDUO (adapalene/benzoyl peroxide)<br>INOVA 4/1 (benzoyl peroxide/salicylic acid)<br>NUOX (benzoyl peroxide/sulfur)<br>PLEXION (sulfacetamide sodium/sulfur)<br>PRASCION (sulfacetamide sodium/sulfur)<br>ROSADERM (sulfacetamide sodium/sulfur)<br>ROSANIL (sulfacetamide sodium/sulfur)<br>ROSULA (sulfacetamide sodium/sulfur)<br>ROSULA (sulfacetamide sodium/sulfur)<br>SULFOXYL (benzoyl peroxide/sulfur)<br>SULFATOL (sulfacetamide<br>sodium/sulfur/urea)<br>VELTIN (clindamycin/tretinoin)<br>ZENCIA WASH (sulfacetamide sodium/sulfur)<br>ZIANA (clindamycin/tretinoin) | present. (In cases of pregnancy, a<br>trial of retinoids will not be<br>required.)<br>In addition, thirty (30) day trials of<br>combinations of the corresponding<br>preferred single agents available<br>are required before non-preferred<br>combination agents will be<br>authorized.  |
| ALZHEIMER'S AGE           | -                |   |   |
|                           |                  | ASE INHIBITORS  |   |
|                           | donepezil        | ARICEPT (donepezil)<br>ARICEPT 23mg (donepezil)<br>ARICEPT ODT(donepezil)<br>COGNEX (tacrine)<br>donepezil ODT<br>EXELON CAPSULE (rivastigmine)<br>EXELON PATCH (rivastigmine)<br>galantamine<br>galantamine ER<br>RAZADYNE (galantamine)<br>RAZADYNE ER (galantamine)<br>rivastigmine  | A thirty (30) day trial of a preferred<br>agent is required before a non-<br>preferred agent in this class will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.<br>Aricept 23mg tablets will be<br>approved when there is a diagnosis<br>of moderate-to-severe Alzheimer's<br>Disease, a trial of donepezil 10mg<br>daily for at least three (3) months,<br>and donepezil 20mg daily for an<br>additional one (1) month.<br>Aricept and donepezil ODT will be<br>approved only when the oral<br>dosage form is not appropriate for<br>the patient. |



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|                           |  |  | Members currently utilizing Exelon patches as of 1/1/2012 may continue.   |
|                           | NMDA RECEPTO   | DR ANTAGONIST  |   |
|                           | NAMENDA (memantine)  |  |   |
|                           | RCOTIC - SHORT ACTING (Non-pai   | contoral) <sup>AP</sup>  |   |
|                           | ASA/codeine<br>codeine<br>dihydrocodeine/ APAP/caffeine<br>hydrocodone/APAP<br>hydrocodone/ibuprofen<br>hydromorphone<br>levorphanol<br>morphine           | ACTIQ (fentanyl)<br>butalbital/APAP/caffeine/codeine<br>butalbital/ASA/caffeine/codeine<br>butorphanol<br>COMBUNOX (oxycodone/ibuprofen)<br>DEMEROL (meperidine)<br>DILAUDID (hydromorphone)<br>fentanyl   | chemically distinct preferred agents<br>(based on narcotic ingredient only),<br>including the generic formulation of<br>a requested non-preferred product,<br>are required before a non-preferred<br>agent will be authorized unless one<br>of the exceptions on the PA form is<br>present.   |
|                           | oxycodone<br>oxycodone/APAP<br>oxycodone/ASA<br>pentazocine/APAP<br>pentazocine/naloxone<br>ROXICET (oxycodone/acetaminophen)<br>tramadol<br>tramadol/APAP | FENTORA (fentanyl)<br>FIORICET W/ CODEINE<br>(butalbital/APAP/caffeine/codeine)<br>FIORINAL W/ CODEINE<br>(butalbital/ASA/caffeine/codeine)<br>LAZANDA (fentanyl)<br>LORCET (hydrocodone/APAP)<br>LORTAB (hydrocodone/APAP)<br>MAGNACET (oxycodone/APAP)<br>meperidine<br>NUCYNTA (tapentadol)<br>OPANA (oxymorphone)<br>ONSOLIS (fentanyl)<br>oxycodone/ibuprofen<br>OXECTA (oxycodone)<br>OXYFAST (oxycodone)<br>OXYFAST (oxycodone)<br>OXYIR (oxycodone)<br>PANLOR (dihydrocodeine/ APAP/caffeine)<br>PERCODAN (oxycodone/ASA)<br>ROXANOL (morphine)<br>RYBIX ODT (tramadol)<br>TALACEN (pentazocine/APAP)<br>TALWIN NX (pentazocine/APAP)<br>REZIX (dihydrocodeine/ APAP/caffeine) <sup>NR</sup> | Fentanyl lozenges and Onsolis will<br>only be approved for a diagnosis of<br>cancer and as an adjunct to a long-<br>acting agent. Neither will be<br>approved for monotherapy.<br>Limits: Unless the patient has<br>escalating cancer pain or another<br>diagnosis supporting increased<br>quantities of short-acting opioids, all<br>short acting solid forms of the<br>narcotic analgesics are limited to<br>120 tablets per 30 days for the<br>purpose of maximizing the use of<br>longer acting medications to<br>prevent unnecessary breakthrough<br>pain in chronic pain therapy. |



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|                           |  | TYLENOL W/CODEINE (APAP/codeine)<br>ULTRACET (tramadol/APAP)<br>ULTRAM (tramadol)<br>VICODIN (hydrocodone/APAP)<br>VICOPROFEN (hydrocodone/ibuprofen)<br>VOPAC (codeine/acetaminophen)<br>XODOL (hydrocodone/acetaminophen)<br>ZAMICET (hydrocodone/APAP)<br>ZYDONE (hydrocodone/acetaminophen)<br>XOLOX (oxycodone/APAP)  |  |
| ANALGESICS, NAF           | RCOTIC - LONG ACTING (Non-pare   | •  |  |
|                           | fentanyl transdermal<br>KADIAN (morphine) 10mg, 20mg, 30mg,<br>50mg, 60mg, 100mg<br>methadone<br>morphine ER tablets<br>OPANA ER (oxymorphone) | AVINZA (morphine)<br>BUTRANS (buprenorphine)<br>CONZIP ER (tramadol)<br>DOLOPHINE (methadone)<br>DURAGESIC (fentanyl)<br>EXALGO ER (hydromorphone)<br>EMBEDA (morphine/naltrexone)<br>KADIAN (morphine) 80mg, 200mg<br>morphine ER capsules<br>MS CONTIN (morphine)<br>NUCYNTA ER (tapentadol)<br>ORAMORPH SR (morphine)<br>oxycodone ER<br>OXYCONTIN (oxycodone)<br>oxymorphone ER<br>RYZOLT ER (tramadol)<br>tramadol ER<br>ULTRAM ER (tramadol) | <ul> <li>Six (6) day trials each of two preferred unique long acting chemical entities are required before a non-preferred agent will be approved unless one of the exceptions on the PDL form is present. The generic form of the requested non-preferred agent, if available, must be tried before the non-preferred agent will be approved.</li> <li>Butrans will be approved if the following criteria are met: <ol> <li>Diagnosis of moderate to severe chronic pain requiring continuous around-the-clock analgesia and</li> <li>Patient cannot take oral medications and has a diagnosis of chronic pain and</li> <li>Needs analgesic medication for an extended period of time and</li> <li>Has had a previous trial** of a non-opioid analgesic medication and</li> <li>Current total daily opioid dose is ≤ 80 mg morphine equivalents daily or dose of transdermal fentanyl is ≤ 12.5 mcg/hr and</li> </ol> </li> </ul> |



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|                           |   |  | <ul> <li>7. Patient is not currently being treated with buprenorphine.</li> <li>**Requirement is waived for patients who cannot swallow</li> <li>Dose optimization is required for achieving equivalent doses of Kadian 80mg and 200mg. AP does not apply.</li> <li>Exception: Oxycodone ER will be authorized if a diagnosis of cancer is submitted without a trial of the preferred agents.</li> </ul>   |
| ANALGESICS (Top           | ical) <sup>AP</sup>   |  |  |
|                           | capsaicin<br>lidocaine<br>lidocaine/prilocaine<br>xylocaine | EMLA (lidocaine/prilocaine)<br>FLECTOR PATCH (diclofenac)<br>LIDODERM PATCH (lidocaine)<br>LIDAMANTLE (lidocaine)<br>LIDAMANTLE HC (lidocaine/hydrocortisone)<br>LMX 4 (lidocaine)<br>PENNSAID (diclofenac)<br>SYNERA (lidocaine/tetracaine)<br>VOLTAREN GEL (diclofenac)<br>ZOSTRIX (capsaicin) | Ten (10) day trials of each of the<br>preferred topical anesthetics<br>(lidocaine, lidocaine/prilocaine, and<br>xylocaine) are required before a<br>non-preferred topical anesthetic will<br>be approved unless one of the<br>exceptions on the PA form is<br>present.<br>Lidoderm patches will be approved<br>for a diagnosis of post-herpetic<br>neuralgia.<br>Thirty (30) day trials of each of the<br>preferred oral NSAIDS and<br>capsaicin are required before<br>Voltaren Gel will be approved<br>unless one of the exceptions on the<br>PA form is present.<br>Flector patches will be approved<br>only for a diagnosis of acute strain,<br>sprain or injury after a five (5) day<br>trial of one of the preferred oral<br>NSAIDs and for a maximum<br>duration of 14 days unless one of |



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|                           |  |   | the exceptions on the PA form is present.  |
| ANDROGENIC AGE            | INTS   |   |  |
|                           | ANDRODERM (testosterone)<br>ANDROGEL (testosterone)  | AXIRON (testosterone)<br>FORTESTA (testosterone)<br>TESTIM (testosterone)   | The non-preferred agents will be<br>approved only if one of the<br>exceptions on the PA form is<br>present.  |
| ANGIOTENSIN MOI           |  |   |  |
|                           |  | IIBITORS  |  |
|                           | benazepril<br>captopril<br>enalapril<br>fosinopril<br>lisinopril<br>quinapril<br>ramipril  | ACCUPRIL (quinapril)<br>ACEON (perindopril)<br>ALTACE (ramipril)<br>CAPOTEN (captopril)<br>LOTENSIN (benazepril)<br>MAVIK (trandolapril)<br>moexipril<br>MONOPRIL (fosinopril)<br>perindopril<br>PRINIVIL (lisinopril)<br>trandolapril<br>UNIVASC (moexipril)<br>VASOTEC (enalapril)<br>ZESTRIL (lisinopril)  | Fourteen (14) day trials of each of<br>the preferred agents in the<br>corresponding group, with the<br>exception of the Direct Renin<br>Inhibitors, are required before a<br>non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present. |
|                           | ACE INHIBITOR CO   | MBINATION DRUGS   |  |
|                           | benazepril/amlodipine<br>benazepril/HCTZ<br>captopril/HCTZ<br>enalapril/HCTZ<br>fosinopril/HCTZ<br>lisinopril/HCTZ<br>quinapril/HCTZ | ACCURETIC (quinapril/HCTZ)<br>CAPOZIDE (captopril/HCTZ)<br>LEXXEL (enalapril/felodipine)<br>LOTENSIN HCT (benazepril/HCTZ)<br>LOTREL (benazepril/amlodipine)<br>moexipril/HCTZ<br>PRINZIDE (lisinopril/HCTZ)<br>TARKA (trandolapril/verapamil)<br>trandolapril/verapamil<br>UNIRETIC (moexipril/HCTZ)<br>VASERETIC (enalapril/HCTZ)<br>ZESTORETIC (lisinopril/HCTZ) |  |
|                           | ANGIOTENSIN II RECEP   | TOR BLOCKERS (ARBs)   |  |



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|                           | AVAPRO (irbesartan)<br>BENICAR (olmesartan)<br>DIOVAN (valsartan)<br>Iosartan<br>MICARDIS (telmisartan)  | ATACAND (candesartan)<br>COZAAR (losartan)<br>EDARBI (azilsartan)<br>eprosartan<br>TEVETEN (eprosartan)   |  |
|                           | ARB COME   | BINATIONS   |  |
|                           | AVALIDE (irbesartan/HCTZ)<br>BENICAR-HCT (olmesartan/HCTZ)<br>DIOVAN-HCT (valsartan/HCTZ)<br>EXFORGE (valsartan/amlodipine)<br>EXFORGE HCT (valsartan/amlodipine/HCTZ)<br>losartan/HCTZ<br>MICARDIS-HCT (telmisartan/HCTZ)               | ATACAND-HCT (candesartan/HCTZ)<br>AZOR (olmesartan/amlodipine)<br>EDARBYCLOR (azilsartan/chlorthalidone) <sup>NR</sup><br>HYZAAR (losartan/HCTZ)<br>TEVETEN-HCT (eprosartan/HCTZ)<br>TRIBENZOR (olmesartan/amlodipine/HCTZ)<br>TWYNSTA (telmisartan/amlodipine) |  |
|                           | DIRECT RENI  | N INHIBITORS  |  |
|                           | AMTURNIDE (aliskiren/amlodipine/HCTZ) <sup>AP</sup><br>TEKAMLO (aliskiren/amlodipine) <sup>AP</sup><br>TEKTURNA (aliskiren) <sup>AP</sup><br>TEKTURNA HCT (aliskiren/HCTZ) <sup>AP</sup><br>VALTURNA (aliskiren/valsartan) <sup>AP</sup> |   | A thirty (30) day trial of one<br>preferred ACE, ARB, or<br>combination agents, at the<br>maximum tolerable dose, is<br>required before Tekturna will be<br>approved.<br>Tekturna HCT, Valturna, Tekamlo<br>or Amturnide will be approved if the<br>criteria for Tekturna are met and the<br>patient also needs the other agents<br>in the combination.  |
| ANTIBIOTICS, GI           |  |   |  |
|                           | ALINIA (nitazoxanide)<br>NEO-FRADIN (neomycin)<br>neomycin<br>metronidazole tablet<br>TINDAMAX (tinidazole)  | DIFICID (fidaxomicin)<br>FLAGYL (metronidazole)<br>FLAGYL ER (metronidazole ER)<br>metronidazole capsule<br>VANCOCIN (vancomycin)<br>XIFAXIN (rifaximin)  | A fourteen (14) day trial of a<br>corresponding generic preferred<br>agent is required before a non-<br>preferred brand agent will be<br>approved.<br>Dificid will be approved if 1) there is<br>a diagnosis of severe <i>C. difficile</i><br>infection and 2) there is no<br>response to prior treatment with<br>vancomycin for 10-14 days.<br>Xifaxin 200 mg will be approved for<br>traveller's diarrhea if 1) there is a |



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|                           |  |  | diagnosis of <i>E. coli</i> diarrhea, 2)<br>patient is between 12 and 18 years<br>old or is 18 years or older and has<br>failed a ten (10) day trial of<br>ciprofloxacin.   |
|                           |  |  | Xifaxin 550 mg will be approved for<br>hepatic encephalopathy if 1) there<br>is a diagnosis of hepatic<br>encephalopathy, 2) patient is 18<br>years or older, and 3) patient has a<br>history of and current treatment with<br>lactulose. |
|                           |  |  | Vancocin will be approved after a fourteen (14) day trial of metronidazole for <i>C. difficile</i> infections of mild to moderate severity unless one of the exceptions on the PA form is present.  |
|                           |  |  | Vancocin will be approved for severe <i>C. difficile</i> infections with no previous trial of metronidazole.  |
| ANTIBIOTICS, INH          | ALED   |  |   |
|                           | TOBI (tobramycin)  | CAYSTON (aztreonam)                                | A 28-day trial of the preferred agent<br>is required before the non-preferred<br>agent will be approved unless one<br>of the exceptions on the PA form is<br>present.   |
| ANTICOAGULANTS            | _  |  |   |
|                           |  | TABLE  | Triple of each of the proferred   |
|                           | ARIXTRA (fondaparinux)<br>FRAGMIN (dalteparin)<br>LOVENOX (enoxaparin) | enoxaparin<br>fondaparinux<br>INNOHEP (tinzaparin) | Trials of each of the preferred<br>agents will be required before a<br>non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present.  |
|                           | OR   | AL   |   |



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|                           | PRADAXA (dabigatran) <sup>AP</sup><br>warfarin<br>XARELTO (rivaroxaban)   |  | Pradaxa will be approved for the<br>diagnosis of non-valvular atrial<br>fibrillation.<br>Xarelto will be approved for the<br>diagnosis of non-valvular atrial<br>fibrillation.<br>Xarelto will be approved for DVT<br>prophylaxis if treatment is limited to<br>35 days for hip replacement<br>surgeries or 12 days for knee<br>replacement surgeries.   |
| ANTICONVULSAN             | TS  |  |  |
|                           | ADJU  | /ANTS  |  |
|                           | carbamazepine<br>CARBATROL (carbamazepine)<br>DEPAKOTE SPRINKLE (divalproex)<br>divalproex EC<br>divalproex DR<br>EPITOL (carbamazepine)<br>FELBATOL (felbamate)<br>gabapentin<br>GABITRIL (tiagabine)<br>levetiracetam<br>lamotrigine<br>lamotrigine chewable<br>LYRICA (pregabalin)<br>oxcarbazepine tablets<br>topiramate<br>TRILEPTAL SUSPENSION (oxcarbazepine)<br>valproic acid<br>zonisamide | BANZEL(rufinamide)<br>carbamazepine XR<br>DEPAKENE (valproic acid)<br>DEPAKOTE (divalproex)<br>DEPAKOTE ER (divalproex)<br>EQUETRO (carbamazepine)<br>FANATREX SUSPENSION (gabapentin) <sup>NR</sup><br>felbamate<br>GRALISE (gabapentin)<br>HORIZANT (gabapentin)<br>KEPPRA (levetiracetam)<br>KEPPRA (levetiracetam)<br>LAMICTAL (lamotrigine)<br>LAMICTAL CHEWABLE (lamotrigine)<br>LAMICTAL CHEWABLE (lamotrigine)<br>LAMICTAL CDT (lamotrigine)<br>LAMICTAL XR (lamotrigine)<br>levetiracetam ER<br>NEURONTIN (gabapentin)<br>ONFI (clobazam) <sup>NR</sup><br>SABRIL (vigabatrin)<br>STAVZOR (valproic acid)<br>TEGRETOL (carbamazepine)<br>TEGRETOL XR (carbamazepine)<br>TOPAMAX (topiramate)<br>TRILEPTAL TABLETS (oxcarbazepine)<br>VIMPAT (lacosamide)<br>ZONEGRAN (zonisamide) | A fourteen (14) day trial of one of<br>the preferred agents in the<br>corresponding group is required for<br>treatment naïve patients with a<br>diagnosis of a seizure disorder<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.<br>A thirty (30) day trial of one of the<br>preferred agents in the<br>corresponding group is required for<br>patients with a diagnosis other than<br>seizure disorders unless one of the<br>exceptions on the PA form is<br>present.<br>Non-preferred anticonvulsants will<br>be approved for patients on<br>established therapies with a<br>diagnosis of seizure disorders with<br>no trials of preferred agents<br>required. In situations where AB-<br>rated generic equivalent products<br>are available, "Brand Medically<br>Necessary" must be hand-written |



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|---------------------------|--|--|---|--|--|
|                           |  |  | <ul> <li>by the prescriber on the prescription<br/>in order for the brand name product<br/>to be reimbursed.</li> <li>Requests for Gralise will be<br/>authorized if the following criteria<br/>are met: <ol> <li>Diagnosis of post herpetic<br/>neuralgia</li> <li>Trial of a tricyclic antidepressant<br/>for a least thirty days</li> <li>Trial of gabapentin immediate<br/>release formulation (positive<br/>response without adequate<br/>duration)</li> <li>Request is for once daily dosing<br/>with 1800 mg. maximum daily</li> </ol></li></ul> |  |  |
|                           | BARBITU  | IRATES <sup>AP</sup>   | dosage.   |  |  |
|                           | mephobarbital<br>phenobarbital<br>primidone                        | MEBARAL (mephobarbital)<br>MYSOLINE (primidone)                        |   |  |  |
|                           | BENZODIA   | ZEPINES <sup>AP</sup>  |   |  |  |
|                           | clonazepam<br>DIASTAT (diazepam rectal)<br>diazepam tablets        | diazepam rectal gel<br>KLONOPIN (clonazepam)                           |   |  |  |
|                           |  | TOINS <sup>AP</sup>  |   |  |  |
|                           | DILANTIN INFATABS (phenytoin)<br>PEGANONE (ethotoin)<br>phenytoin  | CEREBYX (fosphenytoin)<br>DILANTIN (phenytoin)<br>PHENYTEK (phenytoin) |   |  |  |
|                           | SUCCIN   | IMIDES   |   |  |  |
|                           | CELONTIN (methsuximide)<br>ethosuximide<br>ZARONTIN (ethosuximide) |  |   |  |  |
| ANTIDEPRESSANT            | ANTIDEPRESSANTS, OTHER   |  |   |  |  |
|                           | SNF  | RIS <sup>AP</sup>  |   |  |  |



# BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID

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|---------------------------|--|--|---|
|                           | CYMBALTA (duloxetine)<br>venlafaxine ER capsules   | EFFEXOR (venlafaxine)<br>EFFEXOR XR (venlafaxine)<br>PRISTIQ (desvenlafaxine)<br>venlafaxine<br>VENLAFAXINE ER Tablets (venlafaxine)   | A six (6) week trial each of a<br>preferred agent and an SSRI is<br>required before a non-preferred<br>agent will be authorized unless one<br>of the exceptions on the PA form is<br>present.   |
|                           | SECOND GENERATIO   | N NON-SSRI, OTHER <sup>AP</sup>  |   |
|                           | bupropion SR<br>bupropion XL<br>mirtazapine<br>SAVELLA (milnacipran) <sup>AP*</sup><br>trazodone | APLENZIN (bupropion hbr)<br>bupropion IR<br>DESYREL (trazodone)<br>EMSAM (selegiline)<br>nefazodone<br>OLEPTRO ER (trazodone)<br>REMERON (mirtazapine)<br>WELLBUTRIN (bupropion)<br>WELLBUTRIN SR (bupropion)<br>WELLBUTRIN SR (bupropion)<br>VIIBRYD (vilazodone hcl) | * Savella will be approved for a<br>diagnosis of fibromyalgia or a<br>previous thirty (30) day trial of a<br>drug that infers fibromyalgia:<br>gabapentin, Cymbalta, Lyrica,<br>amitriptyline or nortriptyline.   |
|                           |  | ED TCAs  |   |
|                           | imipramine hcl   | imipramine pamoate<br>TOFRANIL (imipramine hcl)<br>TOFRANIL PM (imipramine pamoate)  | A twelve (12) week trial of<br>imipramine hcl is required before a<br>non-preferred TCA will be<br>authorized.  |
| ANTIDEPRESSANT            | ſS, SSRIs <sup>₄</sup>   |  |   |
|                           | citalopram<br>fluoxetine<br>fluvoxamine<br>LEXAPRO (escitalopram)<br>paroxetine<br>sertraline    | CELEXA (citalopram)<br>LUVOX (fluvoxamine)<br>LUVOX CR (fluvoxamine)<br>PAXIL (paroxetine)<br>paroxetine ER<br>PEXEVA (paroxetine)<br>PROZAC (fluoxetine)<br>RAPIFLUX (fluoxetine)<br>SARAFEM (fluoxetine)<br>ZOLOFT (sertraline)                                      | Thirty (30) day trials each of two (2)<br>of the preferred agents are required<br>before a non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present. Upon hospital discharge,<br>patients admitted with a primary<br>mental health diagnosis and have<br>been stabilized on a non-preferred<br>SSRI will receive an authorization<br>to continue that drug. |
|                           |  |  |   |
|                           |  | OR BLOCKERS  |   |
|                           | ondansetron<br>ondansetron ODT   | ANZEMET (dolasetron)<br>KYTRIL (granisetron)<br>granisetron  | A 3-day trial of a preferred agent is required before a non-preferred agent will be authorized unless one   |



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|---------------------------|---|--|---|
|                           | CANNA   | GRANISOL (granisetron)<br>SANCUSO (granisetron)<br>ZOFRAN (ondansetron)<br>ZOFRAN ODT (ondansetron)<br>ZUPLENZ (ondansetron)   | of the exceptions on the PA form is<br>present. PA is required for<br>ondansetron when limits are<br>exceeded.  |
|                           | CANNA   | BINOIDS  | Occurrent will be earth arised early for  |
|                           |   | CESAMET (nabilone)<br>dronabinol<br>MARINOL (dronabinol)   | Cesamet will be authorized only for<br>the treatment of nausea and<br>vomiting associated with cancer<br>chemotherapy for patients who<br>have failed to respond adequately<br>to 3-day trials of conventional<br>treatments such as promethazine or<br>ondansetron and are over 18 years<br>of age.<br>Marinol will be authorized only for<br>the treatment of anorexia<br>associated with weight loss in<br>patients with AIDS or cancer and<br>unresponsive to megestrol; or for<br>the prophylaxis of chemotherapy<br>induced nausea and vomiting<br>unresponsive to 3-day trials of<br>ondansetron or promethazine for<br>patients between the ages of 18<br>and 65. |
|                           | SUBSTANCE P   | ANTAGONISTS  |   |
|                           | EMEND (aprepitant)  |  |   |
| ANTIFUNGALS (Or           |   |  |   |
|                           | clotrimazole<br>fluconazole*<br>ketoconazole <sup>CL</sup><br>nystatin<br>terbinafine <sup>CL</sup> | ANCOBON (flucytosine)<br>DIFLUCAN (fluconazole)<br>flucytosine<br>GRIFULVIN V TABLET (griseofulvin)<br>griseofulvin<br>GRIS-PEG (griseofulvin)<br>itraconazole<br>LAMISIL (terbinafine)<br>MYCELEX (clotrimazole)<br>MYCOSTATIN Tablets (nystatin) | Non-preferred agents will be<br>approved only if one of the<br>exceptions on the PA form is<br>present.<br>*PA is required when limits are<br>exceeded.<br>PA is not required for griseofulvin  |



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|---------------------------|---|--|--|
|                           |   | NIZORAL (ketoconazole)<br>NOXAFIL (posaconazole)<br>ORAVIG BUCCAL (miconazole)<br>SPORANOX (itraconazole)<br>VFEND (voriconazole)<br>voriconazole  | suspension for children up to 6<br>years of age for the treatment of<br>tinea capitis.   |
| ANTIFUNGALS (To           | pical) <sup>AP</sup>  |  |  |
|                           | ANTIFU  | JNGALS   |  |
|                           | econazole<br>ketoconazole<br>MENTAX (butenafine)<br>NAFTIN (naftifine)<br>nystatin<br>ANTIFUNGAL/STER | ciclopirox<br>ERTACZO (sertaconazole)<br>EXELDERM (sulconazole)<br>LOPROX (ciclopirox)<br>MYCOSTATIN (nystatin)<br>NIZORAL (ketoconazole)<br>OXISTAT (oxiconazole)<br>PEDIPIROX-4 (ciclopirox) <sup>NR</sup><br>PENLAC (ciclopirox)<br>SPECTAZOLE (econazole)<br>VUSION (miconazole/petrolatum/zinc oxide)<br>XOLEGEL (ketoconazole) | Fourteen (14) day trials of two (2) of<br>the preferred agents are required<br>before one of the non-preferred<br>agents will be authorized unless<br>one of the exceptions on the PA<br>form is present. If a non-preferred<br>shampoo is requested, a fourteen<br>(14) day trial of one preferred<br>product (ketoconazole shampoo) is<br>required.<br>Oxistat cream will be approved for<br>children 12 and under for tinea<br>corporis, tinea cruris, tinea pedis,<br>and tinea (pityriasis) versicolor. |
|                           | clotrimazole/betamethasone<br>nystatin/triamcinolone  | KETOCON PLUS<br>(ketoconazole/hydrocortisone)<br>LOTRISONE (clotrimazole/betamethasone) <sup>AP</sup><br>MYCOLOG (nystatin/triamcinolone) <sup>AP</sup>  |  |
| ANTIHISTAMINES,           | MINIMALLY SEDATING <sup>AP</sup>  |  |  |
|                           | -   | TAMINES  |  |
|                           | ALAVERT (loratadine)<br>cetirizine<br>loratadine<br>TAVIST-ND (loratadine)                            | ALLEGRA (fexofenadine)<br>CLARINEX Tablets (desloratadine)<br>CLARINEX REDITABS (desloratadine)<br>CLARINEX Syrup (desloratadine)<br>CLARITIN (loratadine)<br>fexofenadine (Rx and OTC)<br>levocetirizine<br>XYZAL (levocetirizine)<br>ZYRTEC (Rx and OTC) (cetirizine)<br>ZYRTEC SYRUP (cetirizine)                                 | Thirty (30) day trials of at least two<br>(2) chemically distinct preferred<br>agents (in the age appropriate<br>form), including the generic<br>formulation of a requested non-<br>preferred product, are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.  |



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|---------------------------|--|---|---|
|                           | ANTIHISTAMINE/DECONO   | SESTANT COMBINATIONS  |   |
|                           | ALAVERT-D (loratadine/pseudoephedrine)<br>cetirizine/pseudoephedrine<br>loratadine/pseudoephedrine<br>SEMPREX-D (acrivastine/ pseudoephedrine) | ALLEGRA-D (fexofenadine/<br>pseudoephedrine)<br>CLARINEX-D (desloratadine/<br>pseudoephedrine)<br>CLARITIN-D (loratadine/pseudoephedrine)<br>fexofenadine/ pseudoephedrine (Rx and<br>OTC)<br>ZYRTEC-D (cetirizine/pseudoephedrine)                       |   |
| ANTIMIGRAINE AG           | ENTS, TRIPTANS <sup>AP</sup>   |   |   |
|                           |  | TANS  |   |
|                           | IMITREX NASAL SPRAY(sumatriptan)<br>IMITREX INJECTION (sumatriptan) <sup>CL</sup><br>naratriptan<br>sumatriptan                                | AMERGE (naratriptan)<br>AXERT (almotriptan)<br>FROVA (frovatriptan)<br>IMITREX tablets (sumatriptan)<br>MAXALT (rizatriptan)<br>MAXALT MLT (rizatriptan)<br>RELPAX (eletriptan)<br>sumatriptan nasal spray/injection <sup>*</sup><br>ZOMIG (zolmitriptan) | Three (3) day trials of each unique<br>chemical entity of the preferred<br>agents are required before a non-<br>preferred agent will be approved<br>unless one of the exceptions on the<br>PA form is present. Quantity limits<br>apply for this drug class.<br>*AP does not apply to nasal spray<br>or injectable sumatriptan. |
|                           |  | TREXIMET (sumatriptan/naproxen sodium)  |   |
|                           |  |   |   |
| ANTIPARKINSON'S           | S AGENTS (Oral)  |   |   |
|                           |  | INERGICS  |   |
|                           | benztropine<br>trihexyphenidyl   | COGENTIN (benztropine)  | Patients starting therapy on drugs<br>in this class must show a<br>documented allergy to all of the<br>preferred agents, in the<br>corresponding class, before a non-<br>preferred agent will be authorized.  |
|                           | COMT IN  | HIBITORS  |   |
|                           |  | COMTAN (entacapone)<br>TASMAR (tolcapone)   |   |



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|---------------------------|---|---|--|
|                           | DOPAMINE  | AGONISTS  |  |
|                           | pramipexole<br>ropinirole   | MIRAPEX (pramipexole)<br>MIRAPEX ER (pramipexole)<br>REQUIP (ropinirole)<br>REQUIP XL (ropinirole)  | Mirapex, Mirapex ER, Requip, and<br>Requip XL will be approved for a<br>diagnosis of Parkinsonism with no<br>trials of preferred agents required.  |
|                           |   | KINSON'S AGENTS   |  |
|                           | amantadine <sup>AP</sup><br>bromocriptine<br>carbidopa/levodopa<br>selegiline<br>STALEVO (levodopa/carbidopa/entacapone)  | AZILECT (rasagiline)<br>ELDEPRYL (selegiline)<br>levodopa/carbidopa ODT<br>LODOSYN (carbidopa)<br>PARCOPA (levodopa/carbidopa)<br>SINEMET (levodopa/carbidopa)<br>ZELAPAR (selegiline)  | Amantadine will be approved only for a diagnosis of Parkinsonism.  |
| ANTIPSYCHOTICS            | , ATYPICAL  |   |  |
|                           | SINGLE IN   | IGREDIENT   |  |
|                           | clozapine<br>GEODON (ziprasidone)<br>INVEGA (paliperidone)<br>INVEGA SUSTENNA (paliperidone)*<br>risperidone ODT<br>risperidone solution<br>SEROQUEL (quetiapine) AP (25mg Tablet Only) | ABILIFY (aripiprazole)<br>CLOZARIL (clozapine)<br>FANAPT (iloperidone)<br>FAZACLO (clozapine)<br>LATUDA (lurasidone)<br>olanzapine<br>olanzapine IM*<br>RISPERDAL (risperidone)<br>RISPERDAL CONSTA (risperidone)*<br>RISPERDAL ODT (risperidone)<br>RISPERDAL SOLUTION (risperidone)<br>SAPHRIS (asenapine)<br>SEROQUEL XR (quetiapine)<br>ZYPREXA (olanzapine)<br>ZYPREXA INTRAMUSCULAR (olanzapine)* | A fourteen (14) day trial of a<br>preferred agent is required for<br>treatment naïve patients before a<br>non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present. Upon discharge, a<br>hospitalized patient stabilized on a<br>non-preferred agent may receive<br>authorization to continue this drug<br>for labeled indications and at<br>recommended dosages.<br>Claims for Seroquel 25 mg will be<br>approved:<br>1. for a diagnosis of<br>schizophrenia<br>or<br>2. for a diagnosis of bipolar<br>disorder<br>or<br>3. when prescribed concurrently<br>with other strengths of<br>Seroquel in order to achieve<br>therapeutic treatment levels.<br>Seroquel 25 mg. will not be |



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|---------------------------|-------------------------------------|---|---|
| DRUG CLASS                |                                     |   | <ul> <li>approved for use as a sedative hypnotic.</li> <li>All antipsychotic agents require prior authorization for children up to six (6) years of age.</li> <li>Abilify will be approved for children between the ages of 6-17 for irritability associated with autism. Abilify will be prior authorized for MDD if the following criteria are met: <ol> <li>The patient is at least 18 years of age.</li> <li>Diagnosis of Major Depressive Disorder (MDD),</li> <li>Evidence of trials of appropriate therapeutic duration (30 days), at the maximum tolerable dose, of at least one agent in two of the following classes: SSRI, SNRI or bupropion in conjunction with Seroquel at doses of 150 mg or more</li> <li>Prescribed in conjunction with an SSRI, SNRI, or bupropion 5. The daily dose does not exceed 15 mg.</li> </ol> </li> </ul> |
|                           |                                     |   | *All injectable antipsychotic<br>products require clinical prior<br>authorization.  |
|                           | ATYPICAL ANTIPSYCHO                 |   |   |
|                           |                                     | SYMBYAX (olanzapine/fluoxetine)                     |   |
| <b>ANTIVIRALS (Oral)</b>  |                                     |   |   |
|                           | ANTI H                              | ERPES   |   |
|                           | acyclovir<br>VALTREX (valacyclovir) | famciclovir<br>FAMVIR (famciclovir)<br>valacyclovir | Five (5) day trials each of the preferred agents are required before the non-preferred agents will  |



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|---------------------------|---|---|--|
|                           |   | ZOVIRAX (acyclovir)   | be authorized unless one of the<br>exceptions on the PA form is<br>present.  |
|                           | ANTI-INF  | LUENZA  | •  |
|                           | RELENZA (zanamivir)<br>TAMIFLU (oseltamivir)  | FLUMADINE (rimantadine)<br>rimantadine<br>amantadine <sup>AP</sup>  | The anti-influenza agents will be approved only for a diagnosis of influenza.  |
| ANTIVIRALS (Topic         | cal) <sup>₄</sup>   |   |  |
|                           | ABREVA (docosanol)<br>DENAVIR (penciclovir)   | ZOVIRAX (acyclovir)   | Five day trials of each of the<br>preferred agents are required<br>before the non-preferred agent will<br>be approved.   |
| ATOPIC DERMATIT           | -   |   |  |
|                           | ELIDEL (pimecrolimus) <sup>AP</sup>   | PROTOPIC (tacrolimus)   | A thirty (30) day trial of a preferred<br>medium or high potency topical<br>corticosteroid is required before<br>coverage of Elidel will be<br>considered; additionally, a thirty<br>(30) day trial of Elidel is required<br>before Protopic will be considered,<br>unless one of the exceptions on the<br>PA form is present. |
| BETA BLOCKERS             | (Oral) & MISCELLANEOUS ANTIAN   | NGINALS (Oral) <sup>₄</sup> P   |  |
|                           | BETA BL   | OCKERS  |  |
|                           | acebutolol<br>atenolol<br>betaxolol<br>bisoprolol<br>metoprolol ER<br>nadolol<br>pindolol<br>propranolol ER<br>sotalol<br>timolol | BETAPACE (sotalol)<br>BLOCADREN (timolol)<br>BYSTOLIC (nebivolol)<br>CARTROL (carteolol)<br>CORGARD (nadolol)<br>INDERAL LA (propranolol)<br>INNOPRAN XL (propranolol)<br>KERLONE (betaxolol)<br>LEVATOL (penbutolol)<br>LOPRESSOR (metoprolol)<br>SECTRAL (acebutolol)<br>TENORMIN (atenolol)<br>TOPROL XL (metoprolol)<br>ZEBETA (bisoprolol) | Fourteen (14) day trials each of<br>three (3) chemically distinct<br>preferred agents, including the<br>generic formulation of a requested<br>non-preferred product, are required<br>before one of the non-preferred<br>agents will be approved unless one<br>of the exceptions on the PA form is<br>present.                  |



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|                 | PREFERRED AGENTS   | NON-PREFERRED AGENTS  | PA CRITERIA  |
|-----------------|--|---|--|
| DRUG CLASS      |  |   |  |
|                 |  |   |  |
|                 | atenolol/chlorthalidone<br>bisoprolol/HCTZ<br>metoprolol/HCTZ<br>nadolol/bendroflumethiazide<br>propranolol/HCTZ | CORZIDE (nadolol/bendroflumethiazide)<br>DUTOPROL (metoprolol ER/HCTZ ER) <sup>NR</sup><br>INDERIDE (propranolol/HCTZ)<br>LOPRESSOR HCT (metoprolol/HCTZ)<br>TENORETIC (atenolol/chlorthalidone)<br>ZIAC (bisoprolol/HCTZ)                  |  |
|                 | BETA- AND ALF  | PHA-BLOCKERS  |  |
|                 | carvedilol<br>labetalol  | COREG (carvedilol)<br>COREG CR (carvedilol)<br>TRANDATE (labetalol)   |  |
|                 | ANTIAN   | GINALS  |  |
|                 |  | RANEXA (ranolazine) <sup>AP</sup>   | Ranexa will be approved for<br>patients with angina who are also<br>taking a calcium channel blocker, a<br>beta blocker, or a nitrite as single<br>agents or a combination agent<br>containing one of these ingredients. |
| BLADDER RELAXA  | ANT PREPARATIONS <sup>AP</sup>   |   |  |
|                 | oxybutynin<br>oxybutynin ER<br>TOVIAZ (fesoterodine)<br>VESICARE (solifenacin)                                   | ENABLEX (darifenacin)<br>DETROL (tolterodine)<br>DETROL LA (tolterodine)<br>DITROPAN (oxybutynin)<br>DITROPAN XL (oxybutynin)<br>GELNIQUE (oxybutynin)<br>OXYTROL (oxybutynin)<br>SANCTURA (trospium)<br>SANCTURA XR (trospium)<br>trospium | A thirty (30) day trial each of the chemically distinct preferred agents is required before a non-preferred agent will be authorized unless one of the exceptions on the PA form is present.                             |
| BONE RESORPTIO  | N SUPPRESSION AND RELATED  |   |  |
| BISPHOSPHONATES |  |   |  |
|                 | alendronate<br>FOSAMAX SOLUTION (alendronate)  | ACTONEL (risedronate)<br>ACTONEL WITH CALCIUM (risedronate/<br>calcium)<br>ATELVIA (risedronate)<br>BONIVA (ibandronate)<br>DIDRONEL (etidronate)<br>FOSAMAX TABLETS (alendronate)<br>FOSAMAX PLUS D (alendronate/vitamin D)                | A 30-day trial of the preferred agent<br>is required before a non-preferred<br>agent will be approved.   |



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|-------------------------------------|---|---|---|
|                                     |   | ZOMETA (zoledronic acid)  |   |
|                                     | OTHER BONE RESORPTION SUPP  | PRESSION AND RELATED AGENTS   |   |
|                                     | MIACALCIN (calcitonin)  | calcitonin<br>EVISTA (raloxifene)<br>FORTEO (teriparatide)<br>FORTICAL (calcitonin)   | Evista will be approved for<br>postmenopausal women with<br>osteoporosis or at high risk for<br>invasive breast cancer.   |
| BPH AGENTS <sup>AP</sup>            |   |   |   |
|                                     | 5-ALPHA-REDUCTA   | SE (5AR) INHIBITORS   |   |
|                                     | AVODART (dutasteride)<br>finasteride  | PROSCAR (finasteride)   | Thirty (30) day trials each of at least<br>two (2) chemically distinct preferred<br>agents, including the generic<br>formulation of a requested non-<br>preferred agent, are required before<br>a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present. |
|                                     | ALPHA B   | LOCKERS   |   |
|                                     | doxazosin<br>tamsulosin<br>terazosin  | alfuzosin<br>CARDURA (doxazosin)<br>CARDURA XL (doxazosin)<br>FLOMAX (tamsulosin)<br>HYTRIN (terazosin)<br>RAPAFLO (silodosin)<br>UROXATRAL (alfuzosin) |   |
|                                     | 5-ALPHA-REDUCTASE (5AR) INHIBIT   | ORS/ALPHA BLOCKER COMBINATION   |   |
|                                     |   | JALYN (dutasteride/tamsulosin)  | Thirty (30) day trials of dutasteride<br>and tamsulosin concurrently are<br>required before the non-preferred<br>agent will be approved.  |
| BRONCHODILATORS & RESPIRATORY DRUGS |   |   |   |
|                                     | ANTICHOL<br>ATROVENT HFA (ipratropium)<br>ipratropium<br>SPIRIVA (tiotropium) | -INERGIC <sup>AP</sup>  | Thirty (30) day trials each of the<br>preferred agents in the<br>corresponding group are required<br>before a non-preferred agent will be   |



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|---------------------------|-----------------------------------|--|---|
|                           |                                   |  | authorized unless one of the exceptions on the PA form is present.  |
|                           | ANTICHOLINERGIC-BETA              | AGONIST COMBINATIONS <sup>AP</sup>   |   |
|                           | COMBIVENT (albuterol/ipratropium) | albuterol/ipratropium<br>DUONEB (albuterol/ipratropium)  | For severely compromised patients,<br>albuterol/ipratropium will be<br>approved if the combined volume of<br>albuterol and ipratropium nebules is<br>inhibitory.  |
|                           | PDE4 IN                           | HIBITOR  |   |
|                           |                                   | DALIRESP (roflumilast)   | Daliresp will be approved when the<br>following criteria are met:<br>1. Patient is ≥ forty (40) years of<br>age and<br>2. Diagnosis of severe chronic<br>obstructive pulmonary disease<br>(COPD) associated with chronic<br>bronchitis and multiple<br>exacerbations requiring systemic<br>glucocorticoids in the preceding six<br>(6) months and<br>3. Concurrent therapy with an<br>inhaled corticosteroid and long-<br>acting bronchodilator and evidence<br>of compliance and<br>4. No evidence of moderate to<br>severe liver impairment (Child-Pugh<br>Class B or C) and<br>5. No concurrent use with strong<br>cytochrome P450 inhibitors<br>(rimampicin, phenobarbital,<br>carbamazepine or phenytoin). |
|                           |                                   | · · · · · · · · · · · · · · · · · ·  |   |
|                           |                                   | SOLUTION <sup>AP</sup>   | Thirty (20) dow trials each of the  |
|                           | albuterol 2.5mg/0.5mL             | ACCUNEB (albuterol)**<br>albuterol 0.63mg & 1.25mg/3mL <sup>AP</sup><br>BROVANA (arformoterol)<br>levalbuterol<br>metaproterenol<br>PERFOROMIST (formoterol) | Thirty (30) day trials each of the<br>chemically distinct preferred agents<br>in their corresponding groups are<br>required before a non-preferred<br>agent in that group will be<br>authorized unless one of the   |



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|---------------------------|--|--|---|
|                           |  | XOPENEX (levalbuterol)   | exceptions on the PA form is<br>present.<br>**No PA is required for ACCUNEB<br>for children up to 5 years of age.   |
|                           | INHALERS, LC   | ONG-ACTING <sup>AP</sup>   |   |
|                           | FORADIL (formoterol)<br>SEREVENT (salmeterol)  | ARCAPTA (indacaterol maleate)  | Thirty (30) day trials each of the<br>preferred agents are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.   |
|                           | INHALERS, SH   | ORT-ACTING <sup>AP</sup>   |   |
|                           | MAXAIR (pirbuterol)<br>PROAIR HFA (albuterol)<br>PROVENTIL HFA (albuterol)<br>VENTOLIN HFA (albuterol) | XOPENEX HFA (levalbuterol)   | Xopenex Inhalation Solution will be<br>approved for 12 months for a<br>diagnosis of asthma or COPD for<br>patients on concurrent asthma<br>controller therapy (either oral or<br>inhaled) with documentation of<br>failure on a trial of albuterol or<br>documented intolerance of<br>albuterol, or for concurrent<br>diagnosis of heart disease. |
|                           | OR   | AL <sup>AP</sup>   |   |
|                           | albuterol<br>terbutaline   | metaproterenol<br>VOSPIRE ER (albuterol)   |   |
| CALCIUM CHANNE            | EL BLOCKERS <sup>AP</sup>  |  |   |
|                           | LONG-2   | ACTING   |   |
|                           | amlodipine<br>diltiazem XR, XT<br>felodipine ER<br>nifedipine ER<br>nisoldipine<br>verapamil ER        | ADALAT CC (nifedipine)<br>CALAN SR (verapamil)<br>CARDENE SR (nicardipine)<br>CARDIZEM CD, LA, SR (diltiazem)<br>COVERA-HS (verapamil)<br>DILACOR XR (diltiazem)<br>DYNACIRC CR (isradipine)<br>ISOPTIN SR (verapamil)<br>NORVASC (amlodipine)<br>PLENDIL (felodipine)<br>PROCARDIA XL (nifedipine)<br>SULAR (nisoldipine) | Fourteen (14) day trials each of the<br>preferred agents are required<br>before a non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present.   |



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|---------------------------|---|--|---|
|                           | SHORT   | TIAZAC (diltiazem)<br>VERELAN/VERELAN PM (verapamil)<br>-ACTING  |   |
|                           | diltiazem<br>verapamil  | CALAN (verapamil)<br>CARDENE (nicardipine)<br>CARDIZEM (diltiazem)<br>DYNACIRC (isradipine)<br>isradipine<br>nicardipine<br>nimodipine<br>nifedipine<br>NIMOTOP (nimodipine)<br>PROCARDIA (nifedipine)   |   |
| CEPHALOSPORINS            | S AND RELATED ANTIBIOTICS (O  |  |   |
|                           |   | A-LACTAMASE INHIBITOR COMBINATIONS   |   |
|                           | amoxicillin/clavulanate   | amoxicillin/clavulanate ER<br>AUGMENTIN XR (amoxicillin/clavulanate)<br>MOXATAG (amoxicillin)  | A five (5) day trial of the preferred<br>agent is required before a non-<br>preferred agent is authorized unless<br>one of the exceptions on the PA<br>form is present.         |
|                           | CEPHALC   | DSPORINS   |   |
|                           | cefaclor<br>cefadroxil<br>cefdinir<br>cefditoren<br>cefpodoxime<br>cefprozil<br>cefuroxime<br>cephalexin<br>SPECTRACEF (cefditoren) | CECLOR (cefaclor)<br>CEDAX (ceftibuten)<br>CEFTIN (cefuroxime)<br>CEFZIL (cefprozil)<br>DURICEF (cefadroxil)<br>KEFLEX (cephalexin)<br>OMNICEF (cefdinir)<br>PANIXINE (cephalexin)<br>RANICLOR (cefaclor)<br>SUPRAX (cefixime)<br>VANTIN (cefpodoxime) |   |
| COLONY STIMULA            |   |  |   |
|                           | LEUKINE (sargramostim)<br>NEUPOGEN (filgrastim)   | NEULASTA (filgrastim)  | A thirty (30) day trial of one of the<br>preferred agents is required before<br>the non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is |



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|---------------------------|---|------------------------------|----------------------------------|
|                           |   |                              | present.                         |
|                           | GENERATION ANTIHISTAMINES                                 |                              |                                  |
|                           |   | , 1 <sup>ST</sup> GENERATION |                                  |
|                           | chlorpheniramine  |                              | See posted list of covered NDCs. |
|                           | clemastine  |                              |                                  |
|                           | diphenhydramine   |                              |                                  |
|                           |   |                              |                                  |
|                           |   |                              |                                  |
|                           | dextromethorphan HBR/promethazine                         |                              | See posted list of covered NDCs. |
|                           | ANTIHISTAMINE-ANTITUSSIVE-D                               | ECONGESTANT COMBINATIONS     |                                  |
|                           | brompheniramine/dextromethorphan                          |                              | See posted list of covered NDCs. |
|                           | HBR/pseudoephedrine<br>chlorpheniramine/dextromethorphan/ |                              |                                  |
|                           | pseudoephedrine   |                              |                                  |
|                           |   |                              |                                  |
|                           |   | NON-NARCOTIC                 | See posted list of sovered NDCs  |
|                           | DELSYM (dextromethorphan polistirex)                      |                              | See posted list of covered NDCs. |
|                           |   | ESTANTS                      |                                  |
|                           | phenylephrine<br>pseudoephedrine                          |                              | See posted list of covered NDCs. |
|                           | pseudoepneume   |                              |                                  |
|                           |   | EXPECTORANTS                 |                                  |
|                           | guaifenesin<br>guaifenesin/dextromethorphan               |                              | See posted list of covered NDCs. |
|                           | •   |                              |                                  |
|                           | pseudoephedrine/chlorpheniramine/                         | ANTICHOLINERGIC COMBINATIONS | See posted list of covered NDCs. |
|                           | scopolamine syrup   |                              |                                  |
|                           | DECONGESTANT-ANTIHIS                                      | STAMINE COMBINATIONS         |                                  |
|                           | phenylephrine HCL/chlorpheniramine maleate                |                              | See posted list of covered NDCs. |
|                           | syrup/drops<br>phenylephrine HCL/promethazine syrup       |                              |                                  |
|                           |   |                              |                                  |



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|---------------------------|--|--|--|
| DRUG CLASS                |  |  |  |
|                           |  |  |  |
|                           | NARCOTIC ANTITUSSIVE-E/                    | CPECTORANT COMBINATION   |  |
| <b>CYTOKINE &amp; CAM</b> | ANTAGONISTS℃                               |  |  |
|                           | ENBREL (etanercept)<br>HUMIRA (adalimumab) | CIMZIA (certolizumab/pegol)<br>KINERET (anakinra)<br>ORENCIA (abatacept) SUBCUTANEOUS<br>SIMPONI (golimumab) | Thirty (30) day trials of each of the preferred agents are required before a non-preferred agent will be approved.   |
|                           |  |  | See additional criteria for treatment<br>of psoriasis or psoriatic arthritis at<br>http://www.dhhr.wv.gov/bms/Pharm<br>acy/Pages/pac.aspx  |
| ERYTHROPOIESIS            |  |  |  |
|                           | PROCRIT (rHuEPO)                           | ARANESP (darbepoetin)<br>EPOGEN (rHuEPO)   | <ul> <li>A thirty (30) day trial of the preferred agent is required before a non-preferred agent will be approved.</li> <li>No evidence of untreated GI bleeding, hemolysis, or Vitamin B-12, iron or folate deficiency.</li> <li>Prior authorization will be given for the erythropoesis agents if the following criteria are met:</li> <li>1. Hemoglobin or Hematocrit less than 10/30 respectively. For renewal, hemoglobin or hematocrit levels greater than 12/36 will require dosage reduction or discontinuation. Exceptions will considered on an individual basis after medical documentation is reviewed.</li> <li>(Laboratory values must be dated within six (6) weeks of request.)</li> <li>2. Transferrin saturation ≥ 20%, ferritin levels ≥100 mg/ml, or on</li> </ul> |



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|---------------------------|---|---|---|--|--|
|                           |   |   | <ul> <li>concurrent therapeutic iron therapy.<br/>(Laboratory values must be dated within three (3) weeks of request. For re-authorization, transferrin saturation or ferritin levels are not required if the patient has been responsive to the erythropoietin agent.</li> <li>3. For HIV-infected patients, endogenous serum erythropoietin level must be ≤ 500mU/ml to initiate therapy.</li> <li>4. No evidence of untreated GI bleeding, hemolysis, or Vitamin B-12, iron or folate deficiency.</li> </ul> |  |  |
| FLUOROQUINOLO             | · · · ·   |   |   |  |  |
|                           | CIPRO (ciprofloxacin) Suspension<br>ciprofloxacin<br>ciprofloxacin ER<br>levofloxacin | AVELOX (moxifloxacin)<br>CIPRO (ciprofloxacin) Tablets<br>CIPRO XR (ciprofloxacin)<br>FACTIVE (gemifloxacin)<br>FLOXIN (ofloxacin)<br>LEVAQUIN (levofloxacin)<br>NOROXIN (norfloxacin)<br>ofloxacin<br>PROQUIN XR (ciprofloxacin) | A five (5) day trial of one of the<br>preferred agents is required before<br>a non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present.  |  |  |
| GENITAL WARTS A           | AGENTS  |   |   |  |  |
|                           | ALDARA (imiquimod)  | CONDYLOX (podofilox)<br>imiquimod<br>podofilox<br>VEREGEN (sinecatechins)<br>ZYCLARA (imiquimod)  | A thirty (30) day trial of the<br>preferred agent is required before a<br>non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.<br>Zyclara will be approved for a<br>diagnosis of actinic keratosis.  |  |  |
| GLUCOCORTICOID            | S (Inhaled) <sup>₄⊳</sup>   |   |   |  |  |
|                           | GLUCOCO   | ORTICOIDS   |   |  |  |
|                           |   |   |   |  |  |



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|----------------|---|--|---|
| DRUG CLASS     | FREI ERRED AGENTS   | NON-FREI ERRED AGENTS  |   |
|                | AEROBID (flunisolide)<br>AEROBID-M (flunisolide)<br>ASMANEX (mometasone)<br>FLOVENT HFA (fluticasone)<br>FLOVENT Diskus (fluticasone)<br>QVAR (beclomethasone)  | ALVESCO (ciclesonide)<br>budesonide<br>PULMICORT (budesonide)*   | Thirty (30) day trials each of the<br>preferred agents are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.<br>Pulmicort Respules do not require a<br>prior authorization for children<br>through 8 years of age or for<br>individuals unable to use an MDI.<br>When children who have been<br>stabilized on Pulmicort Respules<br>reach age 9, prescriptions for the<br>Pulmicort inhaler will be authorized<br>for them.<br>*For children less than 9 years of<br>age and for those who meet the PA<br>requirements, brand Pulmicort is |
|                |   |  | preferred over the generic.   |
|                |   | IODILATOR COMBINATIONS   |   |
|                | ADVAIR (fluticasone/salmeterol)<br>ADVAIR HFA (fluticasone/salmeterol)<br>DULERA (mometasone/formoterol)<br>SYMBICORT(budesonide/formoterol)  |  |   |
| GLUCOCORTICOID | DS (Topical)  |  |   |
|                | VERY HIGH & H   |  |   |
|                | betamethasone dipropionate cream/ointment<br>betamethasone dipropionate/propylene glycol<br>betamethasone valerate ointment<br>clobetasol propionate<br>cream/gel/ointment/solution<br>clobetasol propionate/emollient<br>desoximetasone cream/gel/ointment<br>fluocinonide<br>halobetasol propionate<br>triamcinolone acetonide 0.5% | amcinonide<br>APEXICON (diflorasone diacetate)<br>APEXICON E (diflorasone diacetate)<br>betamethasone dipropionate gel<br>clobetasol propionate foam, lotion, shampoo<br>CLOBEX (clobetasol propionate)<br>CORMAX (clobetasol propionate)<br>diflorasone diacetate<br>diflorasone diacetate/emollient<br>DIPROLENE (betamethasone<br>dipropionate/propylene glycol)<br>DIPROLENE AF (betamethasone<br>dipropionate/propylene glycol) | Five (5) day trials of one (1) form of<br>each preferred unique active<br>ingredient in the corresponding<br>potency group are required before a<br>non-preferred agent will be<br>approved.  |



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|---------------------------|---|--|-------------|
|                           |   | DIPROSONE (betamethasone dipropionate)<br>fluocinonide/emollient<br>halcinonide<br>HALOG (halcinonide)<br>KENALOG 0.5% (triamcinolone acetonide)<br>LIDEX (fluocinonide)<br>LIDEX-E (fluocinonide)<br>LUXIQ (betamethasone valerate)<br>OLUX (clobetasol propionate)<br>OLUX-E (clobetasol propionate/emollient)<br>PSORCON (diflorasone diacetate)<br>TEMOVATE (clobetasol propionate)<br>TEMOVATE.E (clobetasol propionate)<br>TEMOVATE-E (clobetasol<br>propionate/emollient)<br>TOPICORT (desoximetasone)<br>ULTRAVATE (halobetasol propionate)<br>VANOS (fluocinonide)                                      |             |
|                           |   | POTENCY  |             |
|                           | betamethasone dipropionate lotion<br>betamethasone valerate cream<br>desoximetasone 0.05%cream<br>fluocinolone acetonide 0.025%<br>fluticasone propionate<br>hydrocortisone valerate<br>mometasone furoate<br>triamcinolone acetonide 0.025% and 0.1% | ARISTOCORT (triamcinolone)<br>betamethasone valerate lotion<br>BETA-VAL (betamethasone valerate)<br>CLODERM (clocortolone pivalate)<br>CORDRAN/CORDRAN SP (flurandrenolide)<br>CUTIVATE (fluticasone propionate)<br>DERMATOP (prednicarbate)<br>ELOCON (mometasone furoate)<br>hydrocortisone butyrate<br>hydrocortisone butyrate/emollient<br>KENALOG 0.1% (triamcinolone acetonide)<br>LOCOID (hydrocortisone butyrate)<br>LOCOID LIPOCREAM (hydrocortisone<br>butyrate/emollient)<br>prednicarbate<br>TOPICORT LP (desoximetasone)<br>TRIDERM (triamcinolone acetonide)<br>WESTCORT (hydrocortisone valerate) |             |
|                           | LOW PO  | DTENCY   |             |
|                           | desonide<br>fluocinolone acetonide 0.01%<br>hydrocortisone 0.5%, 1%, 2.5%<br>hydrocortisone acetate 0.5%, 1% (Rx & OTC)   | ACLOVATE (alclometasone dipropionate)<br>alclometasone dipropionate<br>CAPEX (fluocinolone acetonide)<br>DERMA-SMOOTHE FS (fluocinolone  |             |



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|---------------------------|---|--|--|
|                           |   | acetonide)<br>DESONATE (desonide)<br>DESOWEN (desonide)<br>LOKARA (desonide)<br>PANDEL (hydrocortisone probutate)<br>VERDESO (desonide)  |  |
| <b>GROWTH HORMO</b>       | NE <sup>cL</sup>  |  |  |
|                           | GENOTROPIN (somatropin)<br>NORDITROPIN NORDIFLEX (somatropin)<br>NORDITROPIN FLEXPRO (somatropin)<br>NUTROPIN AQ NUSPIN (somatropin)                                      | HUMATROPE (somatropin)<br>INCRELEX (mecasermin)<br>NUTROPIN (somatropin)<br>NUTROPIN AQ (somatropin)<br>OMNITROPE (somatropin)<br>SAIZEN (somatropin)<br>SEROSTIM (somatropin)<br>TEV-TROPIN (somatropin)<br>ZORBTIVE (somatropin) | The preferred agents must be tried<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.<br>Patients already on a non-preferred<br>agent will receive authorization to<br>continue therapy on that agent for<br>the duration of the existing PA. |
| H. PYLORI COMBII          | NATION TREATMENTS   |  |  |
|                           | Please use individual components:<br>preferred PPI (Dexilant, omeprazole or<br>pantoprazole)<br>amoxicillin<br>tetracycline<br>metronidazole<br>clarithromycin<br>bismuth | HELIDAC<br>(bismuth/metronidazole/tetracycline)<br>PREVPAC<br>(lansoprazole/amoxicillin/clarithromycin)<br>PYLERA<br>(bismuth/metronidazole/tetracycline)  | A trial of all the individual preferred<br>components (with Dexilant,<br>omeprazole or pantoprazole) at the<br>recommended dosages, frequencies<br>and duration is required before the<br>brand name combination packages<br>will be approved unless one of the<br>exceptions on the PA form is<br>present.    |
| HEPATITIS B TREA          | ATMENTS   |  |  |
|                           | EPIVIR HBV (lamivudine)<br>HEPSERA (adefovir)<br>TYZEKA (telbivudine)   | BARACLUDE (entecavir)  | A thirty (30) day trial of one of the<br>preferred agents is required before<br>the non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.  |
| <b>HEPATITIS C TREA</b>   | ATMENTS <sup>℃</sup>  |  |  |
|                           | INCIVEK (telaprevir) <sup>CL</sup><br>PEGASYS (pegylated interferon)<br>PEG-INTRON (pegylated interferon)   | COPEGUS (ribavirin)<br>INFERGEN (consensus interferon)<br>REBETOL (ribavirin)  | Patients starting therapy in this<br>class must try the preferred agent<br>of a dosage form before a non-  |



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|---------------------------|---|--|---|
|                           | ribavirin<br>VICTRELIS (boceprevir) <sup>CL</sup> | RIBAPAK DOSEPACK (ribavirin)<br>RIBASPHERE (ribavirin) | preferred agent of that dosage form will be authorized.   |
|                           |   |  | See additional criteria for Incivek<br>and Victrelis at<br>http://www.dhhr.wv.gov/bms/Pharm<br>acy/Pages/pac.aspx   |
| HYPERURICEMIA /           | AND GOUT AGENTS                                   |  |   |
|                           | ANTIMI  | ITOTICS  |   |
|                           |   | COLCRYS (colchicine)*                                  | A thirty (30) day trial of one of the<br>preferred agents for the prevention<br>of gouty arthritis attacks<br>(colchicine/probenecid, probenecid,<br>or allopurinol) is required before a<br>non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present.<br>*In the case of acute gouty attacks,<br>a 10-day supply (20 tablets) of<br>Colcrys will be approved per 90<br>days. |
|                           | ANTIMITOTIC-URICO                                 | SURIC COMBINATION                                      |   |
|                           | colchicine/probenecid                             |  |   |
|                           |   | DSURIC   |   |
|                           | probenecid  |  |   |
|                           |   | ASE INHIBITORS   |   |
|                           | allopurinol                                       | ULORIC (febuxostat)<br>ZYLOPRIM (allopurinol)          |   |
| HYPOGLYCEMICS             | INCRETIN MIMETICS/ENHANCER                        | S  |   |
|                           |   | TABLE  |   |
|                           |   |  |   |



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|                           |   | BYDUREON (exenatide)<br>BYETTA (exenatide)<br>SYMLIN (pramlintide)<br>VICTOZA (liraglutide)                                | <ul> <li>Byetta, Symlin, and Victoza will be<br/>subject to the following clinical<br/>edits:</li> <li>Byetta and Victoza will be approved<br/>with a previous history of a thirty<br/>(30) day trial of an oral agent<br/>(sulfonylurea, thiazolindinedione<br/>(TZD) and/ or metformin) and no<br/>evidence of concurrent insulin<br/>therapy.</li> <li>Symlin- History of insulin utilization<br/>in the past 90 days. No gaps in<br/>insulin therapy greater than 30<br/>days.</li> </ul>  |
|                           | OR  | AL <sup>AP</sup>   |  |
|                           | JANUMET (sitagliptin/metformin)<br>JANUVIA (sitagliptin)<br>JUVISYNC (sitagliptin/simvastatin)<br>KOMBIGLYZE XR (saxagliptin/metformin)<br>ONGLYZA (saxagliptin)<br>TRADJENTA (linagliptin) | JANUMET XR (sitagliptin/metformin) <sup>NR</sup><br>JENTADUETO (linagliptin/metformin) <sup>NR</sup>                       | <ul> <li>Januvia/Janumet/Juvisync,<br/>Onglyza/Kombiglyze XR and<br/>Tradjenta will be subject to the<br/>following edits:</li> <li>1. Previous history of a 30-day trial<br/>of metformin.</li> <li>2. Tradjenta will not be approved for<br/>concurrent use with insulin.</li> <li>3. Januvia / Janumet / Juvisync,<br/>Onglyza/Kombiglyze XR will be<br/>approved for concurrent use with<br/>insulin for six (6) month intervals.<br/>For re-authorization, HgBA1C<br/>levels must be less than or equal<br/>(≤) to 7. Current laboratory<br/>values must be submitted.</li> </ul> |
| HYPOGLYCEMICS,            | INSULINS  |  |  |
|                           | HUMALOG (insulin lispro) vials<br>HUMALOG PEN/KWIKPEN (insulin lispro)<br>HUMALOG MIX (insulin lispro/lispro<br>protamine) vials only   | APIDRA (insulin glulisine) <sup>AP</sup><br>HUMALOG MIX PENS (insulin lispro/lispro<br>protamine)<br>HUMULIN PEN (insulin) | To receive Apidra, patients must<br>meet the following criteria:<br>1. be 4 years or older;  |



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|---------------------------|---|---|---|
|                           | HUMULIN (insulin) vials only<br>LANTUS (insulin glargine) all forms<br>LEVEMIR (insulin detemir) all forms<br>NOVOLIN (insulin) all forms<br>NOVOLOG (insulin aspart) all forms<br>NOVOLOG MIX all forms (insulin<br>aspart/aspart protamine) |   | <ol> <li>be currently on a regimen<br/>including a longer-acting or<br/>basal insulin.</li> <li>had a trial of a similar preferred<br/>agent, Novolog or Humalog,<br/>with documentation that the<br/>desired results were not<br/>achieved.</li> </ol> |
| HYPOGLYCEMICS,            |   |   |   |
|                           |   | TINIDES   |   |
|                           | PRANDIN (repaglinide)<br>STARLIX (nateglinide)  | nateglinide   | A thirty (30) day trial of the<br>preferred agent is required before a<br>non-preferred agent will be<br>authorized, unless one of the<br>exceptions on the PA form is<br>present.  |
|                           | MEGLITINIDE C   | COMBINATIONS  |   |
|                           |   | PRANDIMET (repaglinide/metformin)   |   |
| HYPOGLYCEMICS,            | MISCELLANEOUS   |   |   |
|                           | WELCHOL (colesevelam) <sup>AP</sup>   |   | Welchol will be approved for add-on<br>therapy for type 2 diabetes when<br>there is a previous history of a 30-<br>day trial of an oral agent<br>(sulfonylurea, thiazolidinedione<br>(TZD) or metformin).   |
| HYPOGLYCEMICS,            | TZDS  |   |   |
|                           | THIAZOLID   | INEDIONES   |   |
|                           | ACTOS (pioglitazone)  | AVANDIA (rosiglitazone) <sup>AP</sup>   | Treatment naïve patients require a<br>two (2) week trial of Actos before<br>Avandia will be authorized, unless<br>one of the exceptions on the PA<br>form is present.   |
|                           | TZD COME  | BINATIONS   |   |
|                           |   | ACTOPLUS MET (pioglitazone/ metformin)<br>ACTOPLUS MET XR (pioglitazone/<br>metformin)<br>AVANDAMET (rosiglitazone/metformin) <sup>AP</sup> | Patients are required to use the components of Actoplus Met and Duetact separately. Exceptions will be handled on a case-by-case  |



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|---------------------------|---|--|--|
| DRUG CLASS                |   | AVANDARYL (rosiglitazone/glimepiride) <sup>AP</sup><br>DUETACT (pioglitazone/glimepiride)  | basis.   |
| IMPETIGO AGENTS           | S (Topical)   |  |  |
|                           | bacitracin<br>gentamicin sulfate<br>mupirocin                                 | ALTABAX (retapamulin)<br>BACTROBAN (mupirocin)<br>CORTISPORIN (bacitracin/neomycin/<br>polymyxin/HC)   | Ten (10) day trials of at least one<br>preferred agent, including the<br>generic formulation of a requested<br>non-preferred agent, are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.                           |
| INTRANASAL RHIN           | IITIS AGENTS <sup>₄</sup>   |  |  |
|                           | ANTICHO   | LINERGICS  |  |
|                           | ipratropium   | ATROVENT(ipratropium)  | Thirty (30) day trials of the preferred<br>nasal anti-cholinergic, an<br>antihistamine, and corticosteroid<br>groups are required before a non-<br>preferred anti-cholinergic will be<br>approved unless one of the<br>exceptions on the PA form is<br>present.                                |
|                           | ANTIHIS   | TAMINES  |  |
|                           | ASTELIN (azelastine)<br>PATANASE (olopatadine)                                | ASTEPRO (azelastine)<br>azelastine   | Thirty (30) day trials of both<br>preferred intranasal antihistamines<br>and a thirty (30) day trial of one of<br>the preferred intranasal<br>corticosteroids are required before<br>the non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present. |
|                           | CORTICO   | STEROIDS   |  |
|                           | fluticasone propionate<br>NASACORT AQ (triamcinolone)<br>NASONEX (mometasone) | BECONASE AQ (beclomethasone)<br>flunisolide<br>FLONASE (fluticasone propionate)<br>NASALIDE (flunisolide)<br>NASAREL (flunisolide)<br>OMNARIS (ciclesonide)<br>RHINOCORT AQUA (budesonide) | Thirty (30) day trials of each<br>preferred agent in the corticosteroid<br>group are required before a non-<br>preferred corticosteroid agent will<br>be authorized unless one of the<br>exceptions on the PA form is<br>present.  |



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|---------------------------|---|---|---|
|                           |   | triamcinolone<br>VERAMYST (fluticasone furoate)                             |   |
| LEUKOTRIENE MO            | DIFIERS   |   |   |
|                           | ACCOLATE (zafirlukast)<br>SINGULAIR (montelukast) | zafirlukast<br>ZYFLO (zileuton)   | Thirty (30) day trials each of the<br>preferred agents are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.   |
| LIPOTROPICS, OT           | HER (Non-statins) <sup>₄</sup>                    |   |   |
|                           |   | QUESTRANTS  |   |
|                           | colestipol  | COLESTID (colestipol)<br>QUESTRAN (cholestyramine)<br>WELCHOL (colesevelam) | A twelve (12) week trial of one of<br>the preferred agents is required<br>before a non-preferred agent in the<br>corresponding category will be<br>authorized.<br>Welchol will be approved for add-on<br>therapy for type 2 diabetes when<br>there is a previous history of a 30-<br>day trial of an oral agent<br>(sulfonylurea, thiazolidinedione<br>(TZD) or metformin). See<br>HYPOGLYCEMICS,<br>MISCELLANEOUS. |
|                           | CHOLESTEROL ABS                                   | ORPTION INHIBITORS  |   |
|                           |   | ZETIA (ezetimibe)   | Zetia, as monotherapy, will only be<br>approved for patients who cannot<br>take statins or other preferred<br>agents. AP does not apply.<br>Zetia will be approved for add-on<br>therapy only after an insufficient<br>response to the maximum tolerable<br>dose of a statin after 12 weeks of<br>therapy. AP does not apply.   |
|                           | FATTY   | ACIDS   |   |
|                           | FAILT   |   |   |



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|---------------------------|---|--|--|
|                           | LOVAZA (omega-3-acid ethyl esters) <sup>AP</sup>  |  | Lovaza will be approved when the<br>patient is intolerant or not<br>responsive to, or not a candidate for<br>nicotinic acid or fibrate therapy.  |
|                           | FIBRIC ACID   | DERIVATIVES  |  |
|                           | fenofibrate 54mg & 160mg<br>fenofibrate micronized 67mg, 134mg &<br>200mg<br>gemfibrozil<br>TRICOR (fenofibrate nanocrystallized)<br>TRILIPIX (fenofibric acid) | ANTARA (fenofibrate)<br>FENOGLIDE (fenofibrate)<br>FIBRICOR (fenofibric acid)<br>fenofibrate nanocrystallized 145mg<br>LIPOFEN (fenofibrate)<br>LOFIBRA (fenofibrate)<br>LOPID (gemfibrozil)<br>TRIGLIDE (fenofibrate) |  |
|                           | NIA   | CIN  |  |
|                           | niacin<br>NIASPAN (niacin)  | NIACELS (niacin)<br>NIACOR (niacin)<br>NIADELAY (niacin)<br>SLO-NIACIN (niacin)  |  |
| LIPOTROPICS, ST           | ATINS <sup>AP</sup>   |  |  |
|                           | STA   | TINS   |  |
|                           | CRESTOR (rosuvastatin)<br>LESCOL (fluvastatin)<br>LESCOL XL (fluvastatin)<br>LIPITOR (atorvastatin)<br>lovastatin<br>pravastatin<br>simvastatin <sup>CL*</sup>  | ALTOPREV (lovastatin)<br>atorvastatin<br>LIVALO (pitavastatin)<br>MEVACOR (lovastatin)<br>PRAVACHOL (pravastatin)<br>ZOCOR (simvastatin)   | Twelve (12) week trials each of two<br>(2) of the preferred statins,<br>including the generic formulation of<br>a requested non-preferred agent,<br>are required before a non-preferred<br>agent will be authorized unless one<br>of the exceptions on the PA form is<br>present.<br>*Zocor/simvastatin 80mg tablets will<br>require a clinical PA |
|                           |   |  |  |
|                           | ADVICOR (lovastatin/niacin)<br>amlodipine / atorvastatin<br>SIMCOR (simvastatin/niacin ER)  | CADUET (atorvastatin/amlodipine)<br>VYTORIN (simvastatin/ ezetimibe)   | Vytorin will be approved only after<br>an insufficient response to the<br>maximum tolerable dose of Lipitor<br>(atorvastatin) or Crestor<br>(rosuvastatin) after 12 weeks,<br>unless one of the exceptions on the  |



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|---------------------------|---|--|--|
|                           |   |  | PA form is present.<br>*Vytorin 80/10mg tablets will<br>require a clinical PA  |
| MACROLIDES/KET            | OLIDES (Oral)   |  |  |
|                           | KETO  | LIDES  |  |
|                           |   | KETEK (telithromycin)  | Requests for telithromycin will be<br>authorized if there is documentation<br>of the use of any antibiotic within<br>the past 28 days.   |
|                           | MACR  | OLIDES   |  |
|                           | azithromycin<br>clarithromycin<br>erythromycin  | BIAXIN (clarithromycin)<br>BIAXIN XL (clarithromycin)<br>clarithromycin ER<br>E.E.S. (erythromycin ethylsuccinate)<br>E-MYCIN (erythromycin)<br>ERYPED (erythromycin ethylsuccinate)<br>ERY-TAB (erythromycin)<br>ERYTHROCIN (erythromycin stearate)<br>erythromycin estolate<br>PCE (erythromycin)<br>ZITHROMAX (azithromycin)<br>ZMAX (azithromycin) | Five (5) day trials each of the<br>preferred agents are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.   |
| MULTIPLE SCLER            | OSIS AGENTS <sup>CL, AP</sup>   |  |  |
|                           | INTERF  | ERONS  |  |
|                           | AVONEX (interferon beta-1a)<br>BETASERON (interferon beta-1b)<br>REBIF (interferon beta-1a) | EXTAVIA (interferon beta-1b)   | A 30-day trial of a preferred agent<br>will be required before a non-<br>preferred agent will be approved.   |
| NON-INTERFERONS           |   |  |  |
|                           | COPAXONE (glatiramer)   | AMPYRA (dalfampridine)*<br>GILENYA (fingolimod)**<br>TYSABRI (natalizumab)***  | A 30-day trial of the preferred agent<br>will be required before a non-<br>preferred agent will be approved.<br>*Amypra will be prior authorized if<br>the following conditions are met:<br>1. Diagnosis of multiple sclerosis |



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|---------------------------|---|---|--|
|                           |   |   | <ol> <li>No history of seizures</li> <li>No evidence of moderate or<br/>severe renal impairment</li> <li>Initial prescription will be<br/>approved for 30 days only.</li> <li>** Gilenya: PA Criteria</li> <li>A diagnosis of a relapsing form<br/>of multiple sclerosis AND</li> <li>Medication is prescribed by a<br/>neurologist AND</li> <li>History of a thirty (30) trial of one<br/>of the preferred agents for<br/>multiple sclerosis unless <i>one of</i><br/>the exceptions on the PA form is<br/>present AND</li> <li>Dosage is limited to one tablet<br/>per day.<br/>(AP does not apply.)</li> <li>***Tysabri will only be <i>approved</i> for<br/>members who are enrolled in the<br/>TOUCH Prescribing Program. AP<br/>does not apply.</li> </ol> |
| MUSCLE RELAXAN            | NTS (Oral) <sup>₄⊳</sup>                          |   |  |
|                           | ACUTE MUSCULOSKELE                                | TAL RELAXANT AGENTS   |  |
|                           | chlorzoxazone<br>cyclobenzaprine<br>methocarbamol | AMRIX (cyclobenzaprine)<br>carisoprodol<br>carisoprodol/ASA<br>carisoprodol/ASA/codeine<br>cyclobenzaprine ER<br>FEXMID (cyclobenzaprine)<br>FLEXERIL (cyclobenzaprine)<br>LORZONE (chlorzoxazone)<br>metaxalone<br>methocarbamol/ASA<br>orphenadrine<br>orphenadrine/ASA/caffeine<br>PARAFON FORTE DSC (chlorzoxazone) | Thirty (30) day trials of the preferred<br>acute musculoskeletal relaxants are<br>required before a non-preferred<br>acute musculoskeletal agent will be<br>approved, with the exception of<br>carisoprodol.<br>Thirty (30) day trials of the preferred<br>acute musculoskeletal relaxants<br>and Skelaxin are required before<br>carisoprodol will be approved.   |



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|---------------------------|--|--|---|
|                           |  | ROBAXIN (methocarbamol)<br>SKELAXIN (metaxalone)<br>SOMA (carisoprodol)<br>SOMA COMPOUND (carisoprodol /ASA)<br>SOMA COMP w/ COD (carisoprodol/ASA/<br>codeine)  |   |
|                           |  | AGENTS USED FOR SPASTICITY   |   |
|                           | baclofen<br>dantrolene<br>tizanidine <mark>tablets</mark>  | DANTRIUM (dantrolene)<br>tizanidine capsules<br>ZANAFLEX (tizanidine)  | Thirty (30) day trials of the preferred<br>skeletal muscle relaxants<br>associated with the treatment of<br>spasticity (are required before non-<br>preferred agents will be approved<br>unless one of the exceptions on the<br>PA form is present. |
| NSAIDS <sup>AP</sup>      | NON-SE   | LECTIVE  |   |
|                           | diclofenac<br>etodolac<br>fenoprofen<br>flurbiprofen<br>ibuprofen (Rx and OTC)<br>INDOCIN (indomethacin) (suspension only)<br>indomethacin<br>ketorolac<br>naproxen (Rx only)<br>oxaprozin<br>sulindac | ADVIL (ibuprofen)<br>ANAPROX (naproxen)<br>ANSAID (flurbiprofen)<br>CAMBIA (diclofenac)<br>CATAFLAM (diclofenac)<br>CLINORIL (sulindac)<br>DAYPRO (oxaprozin)<br>FELDENE (piroxicam)<br>INDOCIN (indomethacin)<br>ketoprofen<br>Retoprofen ER<br>LODINE (etodolac)<br>meclofenamate<br>mefenamic acid<br>MOTRIN (ibuprofen)<br>nabumetone<br>NALFON (fenoprofen)<br>NAPRELAN (naproxen)<br>NAPROSYN (naproxen)<br>NUPRIN (ibuprofen)<br>ORUDIS (ketoprofen)<br>piroxicam<br>PONSTEL (meclofenamate)<br>SPRIX (ketorolac)<br>tolmetin | Thirty (30) day trials of each of the<br>preferred agents are required<br>before a non-preferred agent will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.  |



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|---------------------------|--|---|--|
|                           |  | VOLTAREN (diclofenac)<br>ZIPSOR (diclofenac potassium)  |  |
|                           | NSAID/GI PROTECT   | ANT COMBINATIONS  |  |
|                           |  | ARTHROTEC (diclofenac/misoprostol)<br>VIMOVO (naproxen/esomeprazole)  |  |
|                           | COX-II SE  | ELECTIVE  |  |
|                           | meloxicam  | CELEBREX (celecoxib)<br>MOBIC (meloxicam)   | <ul> <li>Requests for COX-2 Inhibitor agents will be authorized if the following criteria are met:</li> <li>Agent is requested for treatment of a chronic condition, and</li> <li>a. Patient is greater than or equal to 70 years of age, or</li> <li>b. Patient is currently on anticoagulation therapy, or</li> <li>c. Patient has a history or risk of a serious GI complication.</li> </ul>        |
| OPHTHALMIC ANT            | IBIOTICS (FLUOROQUINOLONES   | & SELECT MACROLIDES) <sup>AP</sup>  |  |
|                           | ciprofloxacin<br>MOXEZA (moxifloxacin)<br>ofloxacin<br>VIGAMOX (moxifloxacin)<br>**The American Academy of Ophthalmology<br>guidelines on treating bacterial conjunctivitis<br>recommend as first line treatment options:<br>erythromycin ointment, sulfacetamide drops,<br>or polymyxin/trimethoprim drops. Alternative<br>treatments include bacitracin ointment,<br>sulfacetamide ointment, polymyxin/bacitracin<br>ointment, fluoroquinolone drops, or<br>azithromycin drops. All generic forms of | AZASITE (azithromycin)<br>BESIVANCE (besifloxacin)<br>CILOXAN (ciprofloxacin)<br>levofloxacin<br>OCUFLOX (ofloxacin)<br>QUIXIN (levofloxacin)<br>ZYMAXID (gatifloxacin) | Five (5) day trials of each of the<br>preferred agents are required<br>before non-preferred agents will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present.<br>**A prior authorization is required<br>for the fluoroquinolone agents for<br>patients under 21 years of age<br>unless there has been a trial of a<br>first line treatment option within the<br>past 10 days. |



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|--|--|---|---|--|
|  | ophthalmic erythromycin, sulfacetamide, and<br>polymyxin/trimethoprim, polymyxin/bacitracin<br>and bacitracin are preferred.                         |   |   |  |
| <b>OPHTHALMIC ANT</b>                              | I-INFLAMMATORIES   |   |   |  |
|  | flurbiprofen<br>ketorolac 0.4%<br>NEVANAC (nepafenac)  | ACULAR LS (ketorolac)<br>ACUVAIL 0.45% (ketorolac tromethamine) <sup>AP</sup><br>BROMDAY (bromfenac)<br>diclofenac <sup>AP</sup><br>DUREZOL (difluprednate) <sup>AP</sup><br>LOTEMAX (loteprednol)<br>VEXOL (rimexolone)<br>XIBROM (bromfenac)  | Five (5) day trials of each of the<br>preferred ophthalmic anti-<br>inflammatory agents are required<br>before non-preferred agents will be<br>authorized unless one of the<br>exceptions on the PA form is<br>present. |  |
| <b>OPHTHALMICS FO</b>                              | R ALLERGIC CONJUNCTIVITIS  |   |   |  |
|  | ALAWAY (ketotifen)<br>ALREX (loteprednol)<br>cromolyn<br>ketorolac 0.5%<br>PATADAY (olopatadine)<br>PATANOL (olopatadine)<br>ZADITOR OTC (ketotifen) | ACULAR (ketorolac)<br>ALAMAST (pemirolast) <sup>AP</sup><br>ALOCRIL (nedocromil) <sup>AP</sup><br>ALOMIDE (lodoxamide) <sup>AP</sup><br>azelastine<br>BEPREVE (bepotastine) <sup>AP</sup><br>CROLOM (cromolyn) <sup>AP</sup><br>DUREZOL (difuprednate) <sup>NR</sup><br>ELESTAT (epinastine) <sup>AP</sup><br>EMADINE (emedastine) <sup>AP</sup><br>epinastine<br>ketotifen<br>LASTACAFT (alcaftadine)<br>OPTICROM (cromolyn) <sup>AP</sup><br>OPTIVAR (azelastine)<br>ZYRTEC ITCHY EYE (ketotifen) <sup>AP</sup> | Thirty (30) day trials of each of<br>three (3) of the preferred agents are<br>required before non-preferred<br>agents will be authorized, unless<br>one of the exceptions on the PA<br>form is present.                 |  |
| OPHTHALMICS, GLAUCOMA AGENTS<br>COMBINATION AGENTS |  |   |   |  |
|  | COMBIGAN (brimonidine/timolol)   | COSOPT (dorzolamide/timolol)  | Authorization for a non-preferred   |  |
|  | dorzolamide/timolol  |   | agent will only be given if there is<br>an allergy to the preferred agents.   |  |
|  |  | OCKERS  |   |  |
|  | betaxolol<br>BETOPTIC S (betaxolol)  | BETAGAN (levobunolol)<br>BETIMOL (timolol)  |   |  |



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|---------------------------|--|---|---|
|                           | carteolol<br>levobunolol<br>metipranolol<br>timolol  | ISTALOL (timolol)<br>OPTIPRANOLOL (metipranolol)<br>TIMOPTIC (timolol)                                  |   |
|                           |  | DRASE INHIBITORS  |   |
|                           | AZOPT (brinzolamide)<br>dorzolamide  | TRUSOPT (dorzolamide)   |   |
|                           | PARASYMPA  | THOMIMETICS   |   |
|                           | CARBOPTIC (carbachol)<br>ISOPTO CARBACHOL (carbachol)<br>PHOSPHOLINE IODIDE (echothiophate<br>iodide)<br>pilocarpine | ISOPTO CARPINE (pilocarpine)<br>PILOPINE HS (pilocarpine)   |   |
|                           | PROSTAGLAN   | IDIN ANALOGS  |   |
|                           | latanoprost<br>LUMIGAN (bimatoprost)<br>TRAVATAN-Z (travoprost)  | XALATAN (latanoprost)<br><mark>ZIOPTAN (tafluprost)</mark>  |   |
|                           |  | OMIMETICS   |   |
|                           | ALPHAGAN P (brimonidine)<br>brimonidine 0.2%<br>dipivefrin   | brimonidine 0.15%<br>PROPINE (dipivefrin)   |   |
| OTIC FLUOROQUII           |  |   |   |
|                           | CIPRODEX (ciprofloxacin/dexamethasone)*<br>ofloxacin   | CIPRO HC (ciprofloxacin/hydrocortisone)<br>CETRAXAL 0.2% SOLUTION (ciprofloxacin)<br>FLOXIN (ofloxacin) | Five (5) day trials of each of the<br>preferred agents are required<br>before a non-preferred agent will be<br>approved unless one of the<br>exceptions on the PA form is<br>present. |
|                           |  |   | *Ciprodex is limited to patients 8<br>years of age and younger. Age<br>exceptions will be handled on a<br>case-by-case basis.   |
| PANCREATIC ENZ            | YMES <sup>ap</sup>   |   |   |



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|---|---|---|--|--|
|   | CREON<br>ZENPEP   | PANCREAZE<br>PANCRELIPASE 5000  | A thirty (30) day trial of a preferred<br>agent is required before a non-<br>preferred agent will be authorized<br>unless one of the exceptions on the<br>PA form is present.<br>Non-preferred agents will be<br>approved for members with cystic  |  |
|   |   |   | fibrosis.  |  |
| PARATHYROID AG                                |   |   |  |  |
|   | calcitriol<br>HECTOROL (doxercalciferol)<br>vitamin d 2 (ergocalciferol) (Rx and OTC)*<br>vitamin d 3 (cholecalciferol) (Rx and OTC)* | DRISDOL (ergocalciferol)<br>ROCALTROL (calcitriol)<br>SENSIPAR (cinacalcet)   | A thirty (30) day trial of a preferred<br>agent will be required before a non-<br>preferred agent will be approved.  |  |
|   | ZEMPLAR (paricalcitol)  |   | *See Covered List  |  |
| PEDICULICIDES/SO                              | CABICIDES (Topical) <sup>₄</sup>  |   |  |  |
|   | OVIDE (malathion)<br>permethrin (Rx and OTC)<br>pyrethrins-piperonyl butoxide   | EURAX (crotamiton)<br>lindane<br>LYCELLE (topical gel)<br>malathion 0.5% lotion<br>NATROBA (spinosad)<br>ULESFIA 5% LOTION (benzyl alcohol)             | Trials of the preferred agents<br>(which are age and weight<br>appropriate) are required before<br>non-preferred agents will be<br>approved unless one of the<br>exceptions on the PA form is<br>present.  |  |
| PHOSPHATE BIND                                | ERS   |   |  |  |
|   | calcium acetate<br>FOSRENOL (lanthanum)<br>RENAGEL (sevelamer)<br>RENVELA (sevelamer carbonate)                                       | ELIPHOS (calcium acetate)<br>PHOSLYRA (calcium acetate)   | Thirty (30) day trials of at least two<br>preferred agents are required<br>unless one of the exceptions on the<br>PA form is present.  |  |
| PLATELET AGGREGATION INHIBITORS <sup>AP</sup> |   |   |  |  |
|   | AGGRENOX (dipyridamole/ASA)<br>cilostazol<br>PLAVIX (clopidogrel)   | BRILINTA (ticagrelor)<br>dipyridamole<br>EFFIENT (prasugrel)<br>PERSANTINE (dipyridamole)<br>PLETAL (cilostazol)<br>TICLID (ticlopidine)<br>ticlopidine | A thirty (30) day trial of a preferred<br>agent is required before a non-<br>preferred agent will be approved<br>unless one of the exceptions on the<br>PA form is present.<br>Effient will be approved for acute<br>coronary syndrome when it is to be<br>managed by acute or delayed<br>percutaneous coronary intervention<br>(PCI). Three (3) day emergency |  |



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|---------------------------|---|--|---|
|                           |   |  | supplies of Effient are available when necessary. |
| PRENATAL VITAM            | INS   |  |   |
|                           | prenatal vitamin 27 w/calcium/ferrous<br>fumarate/folic acid<br>prenatal vitamins 28 w/calcium/iron ps<br>complex/folic acid<br>prenatal vitamins/ferrous<br>fumarate/docusate/folic acid<br>prenatal vitamins/ferrous fumarate/folic acid<br>prenatal vitamins/ferrous fumarate/folic<br>acid/selenium<br>prenatal vitamins/iron, carbonyl/folic acid<br>prenatal vitamin no. 15/iron, carbonyl/folic<br>acid/docusate sod<br>prenatal vitamin no. 16/iron, carbonyl/folic<br>acid/docusate sod<br>prenatal vitamin no. 17/iron, carbonyl/folic<br>acid/docusate sod<br>prenatal vitamin no. 18/iron, carbonyl/folic<br>acid/docusate sod<br>prenatal vitamin w-o calcium/ferrous<br>fumarate/folic acid<br>prenatal vitamin w-o vit a/fe carbonyl-fe<br>fumarate/fa | CARENATAL DHA<br>CITRANATAL DHA<br>COMBI RX<br>FOLBECAL<br>DUET/DUET DHA<br>FOLTABS PLUS DHA<br>NATACHEW<br>NATAFORT<br>NATELLE PLUS W/DHA<br>NEEVO<br>NOVANATAL<br>OB-NATAL ONE<br>OPTINATE<br>PRECARE/PRECARE PREMIER<br>PREMESIS<br>PRENATAL RX<br>PRENATAL RX<br>PRENATAL RX<br>PRENATAL RX<br>PRENATAL U<br>prenatal vitamins/ferrous bis-glycinate<br>chelate/folic acid<br>prenatal vitamins ron. 20/iron bisgly/folic<br>acid/DHA<br>prenatal vitamins no. 22/iron,<br>carbonyl/FA/docusate/DHA<br>prenatal vitamins w-CA, FE, FA (<1 mg)<br>prenatal vitamins w-o CA no. 5/ferrous<br>fumarate/folic acid<br>prenatal vitamins w-o CA no. 5/ferrous<br>fumarate/folic acid<br>prenatal vitamins w-o CA no. 5/ferrous<br>fumarate/folic acid<br>prenatal vitamins w-o CA no. 2<br>prenatal vitamins w-o calcium no. 9/iron/folic<br>acid<br>PRENATE DHA/PRENATE ELITE<br>PRENAXE<br>PRIMACARE<br>RENATE/RENATE DHA | See posted list of covered NDCs.                  |



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|---------------------------|---|---|--|
|                           |   | SELECT-OB<br>TANDEM DHA/TANDEM OB   |  |
| PROTON PUMP INI           | HIBITORS <sup>AP</sup>  |   |  |
|                           | DEXILANT (dexlansoprazole)<br>omeprazole<br>pantoprazole                  | ACIPHEX (rabeprazole)<br>lansoprazole<br>NEXIUM (esomeprazole)<br>NEXIUM PACKETS (esomeprazole)<br>omeprazole/sodium bicarbonate<br>PREVACID capsules (lansoprazole)<br>PREVACID Solu-Tabs (lansoprazole)<br>PRILOSEC (omeprazole)<br>PROTONIX (pantoprazole)<br>ZEGERID OTC (omeprazole) | Sixty (60) day trials of each of the<br>preferred agents, inclusive of a<br>concurrent thirty (30) day trial at the<br>maximum dose of an H₂ antagonist<br>are required before a non-preferred<br>agent will be approved unless one<br>of the exceptions on the PA form is<br>present<br>Prior authorization is not required<br>for Prevacid Solu-Tabs for patients<br>≤8 years of age.  |
| PSORIATIC AGENT           | TS - TOPICAL  |   |  |
|                           | calcipotriene ointment<br>DOVONEX (calcipotriene)<br>TAZORAC (tazarotene) | calcipotriene solution<br>TACLONEX<br>(calcipotriene/betamethasone)<br>VECTICAL (calcitriol)  | Thirty (30) day trials of two (2)<br>preferred unique chemical entities<br>are required before non-preferred<br>agents will be approved unless one<br>of the exceptions on the PA form is<br>present.  |
| PULMONARY ANT             | <b>HYPERTENSIVES - ENDOTHELIN</b>   | RECEPTOR ANTAGONISTS <sup>CL</sup>  |  |
|                           | LETAIRIS (ambrisentan)  | TRACLEER (bosentan)   | Letairis will be approved for the<br>treatment of pulmonary artery<br>hypertension (PAH) World Health<br>Organization (WHO) Group I to<br>improve exercise ability and<br>decrease the rate of clinical<br>deterioration.<br>Tracleer will be approved for the<br>treatment of pulmonary artery<br>hypertension (PAH) (WHO Group I)<br>in patients with World Health<br>Organization (WHO) Class II, III, or<br>IV symptoms to improve exercise<br>capacity and decrease the rate of<br>clinical deterioration AND when<br>there has been a failure with |



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|---------------------------|---|---|--|
|                           |   |   | Letairis.  |
| PULMONARY ANTI            | HYPERTENSIVES – PDE5s <sup>cl</sup>         |   |  |
|                           | ADCIRCA (tadalafil)<br>REVATIO (sildenafil) |   |  |
| PULMONARY ANTI            | HYPERTENSIVES – PROSTACYCI                  | _INS <sup>c∟</sup>  |  |
|                           | epoprostenol<br>VENTAVIS (iloprost)         | FLOLAN (epoprostenol)<br>REMODULIN (treprostinil sodium)<br>TYVASO (treprostinil)   | Ventavis will only be approved for<br>the treatment of pulmonary artery<br>hypertension (WHO Group 1) in<br>patients with NYHA Class III or IV<br>symptoms.<br>Remodulin and Tyvaso will be<br>approved only after a 30-day trial of<br>Ventavis unless one of the<br>exceptions on the PA form is<br>present. |
| SEDATIVE HYPNO            | TICS <sup>AP</sup>                          |   |  |
|                           | BENZODI                                     | AZEPINES  |  |
|                           | temazepam                                   | DALMANE (flurazepam)<br>DORAL (quazepam)<br>estazolam<br>flurazepam<br>HALCION (triazolam)<br>RESTORIL (temazepam)<br>triazolam   | Fourteen (14) day trials of the<br>preferred agents in both categories<br>are required before a non-preferred<br>agent will be authorized unless one<br>of the exceptions on the PA form is<br>present.  |
|                           |   | IERS  |  |
|                           | zolpidem                                    | AMBIEN (zolpidem)<br>AMBIEN CR (zolpidem)<br>chloral hydrate<br>EDLUAR SL (zolpidem)<br>LUNESTA (eszopiclone)<br>ROZEREM (ramelteon)<br>SILENOR (doxepin)<br>SOMNOTE (chloral hydrate)<br>SONATA (zaleplon)<br>zaleplon<br>zolpidem tartrate ER |  |



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|---------------------------|---|--|--|
|                           |   | ZOLPIMIST SPRAY (zolpidem  |  |
| STIMULANTS AND            | RELATED AGENTS  |  |  |
|                           |   | TAMINES  |  |
|                           | amphetamine salt combination<br>dextroamphetamine<br>VYVANSE (lisdexamfetamine)   | ADDERALL (amphetamine salt combination)<br>ADDERALL XR (amphetamine salt<br>combination)<br>amphetamine salt combination ER<br>DESOXYN (methamphetamine)<br>DEXEDRINE (dextroamphetamine)<br>DEXTROSTAT (dextroamphetamine)<br>methamphetamine<br>PROCENTRA (dextroamphetamine) <sup>NR</sup>                              | Members currently utilizing Adderall<br>XR as of 1/1/2012 may continue<br>use until 6/30/2012.<br>Except for Strattera, PA is required<br>for adults >18 years.<br>One of the preferred agents in each<br>group (amphetamines and non-<br>amphetamines) must be tried for<br>thirty (30) days before a non-<br>preferred agent will be authorized.<br>Thirty (30) day trials of at least<br>three<br>(3) antidepressants are required<br>before amphetamines will be<br>approved for depression.<br>Provigil will only be approved for<br>patients >16 years of age with a<br>diagnosis of narcolepsy. |
|                           | NON-AMP   | HETAMINE   |  |
|                           | CONCERTA (methylphenidate)<br>DAYTRANA (methylphenidate)<br>FOCALIN (dexmethylphenidate)<br>FOCALIN XR (dexmethylphenidate)<br>guanfacine<br>INTUNIV (guanfacine extended-release)<br>METADATE CD (methylphenidate)<br>methylphenidate<br>methylphenidate ER (Generic Ritalin SR,<br>Metadate ER, Methylin ER)<br>STRATTERA (atomoxetine) | dexmethylphenidate<br>KAPVAY ER (clonidine)<br>METADATE ER (methylphenidate)<br>methylphenidate ER (Generic Concerta)<br>methylphenidate ER (Generic Ritalin LA)<br>NUVIGIL (armodafinil)<br>pemoline<br>PROVIGIL (modafinil)<br>RITALIN (methylphenidate)<br>RITALIN LA (methylphenidate)<br>RITALIN-SR (methylphenidate) | <ul> <li>Strattera will not be approved for concurrent administration with amphetamines or methylphenidates, except for 30 days or less for tapering purposes.</li> <li>Strattera is limited to a maximum of 100mg per day.</li> <li>Kapvay will be approved if the following criteria are met: <ol> <li>Fourteen (14) day trials of at least one preferred product from the amphetamine and non-amphetamine class and</li> <li>A fourteen (14) day trial of Strattera and</li> </ol> </li> </ul>  |



# BUREAU FOR MEDICAL SERVICES WEST VIRGINIA MEDICAID

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|----------------------------|---|--|---|
|                            |   |  | <ol> <li>A fourteen (14) day trial of<br/>clonidine (for Kapvay) unless<br/>one of the exceptions on the<br/>PA form is present or</li> <li>In cases of a diagnosis of<br/>Tourette's syndrome, tics,<br/>autism or disorders included<br/>in the autism spectrum, only a<br/>fourteen (14) day trial of<br/>clonidine (for Kapvay) is<br/>required for approval.</li> </ol>                                  |
| TETRACYCLINES <sup>A</sup> |   | ADOXA (downeysling manchudrate)  | $\Lambda$ top (10) dow trial of each of the   |
|                            | doxycycline hyclate<br>minocycline capsules<br>tetracycline   | ADOXA (doxycycline monohydrate)<br>demeclocycline*<br>DORYX (doxycycline hyclate)<br>doxycycline hyclate delayed release<br>doxycycline monohydrate<br>DYNACIN (minocycline)<br>MINOCIN (minocycline)<br>minocycline SR capsules<br>minocycline tablets<br>MONODOX (doxycycline monohydrate)<br>ORACEA (doxycycline monohydrate)<br>SOLODYN (minocycline)<br>SUMYCIN (tetracycline)<br>VIBRAMYCIN SYRUP (doxycycline calcium)<br>VIBRAMYCIN (doxycycline hyclate)<br>VIBRAMYCIN (doxycycline monohydrate)<br>VIBRAMYCIN (doxycycline monohydrate)<br>VIBRAMYCIN (doxycycline monohydrate)<br>VIBRAMYCIN (doxycycline monohydrate)<br>VIBRATABS (doxycycline hyclate) | A ten (10) day trial of each of the<br>preferred agents is required before<br>a non-preferred agent will be<br>approved.<br>*Demeclocycline will be approved<br>for conditions caused by<br>susceptible strains of organisms<br>designated in the product<br>information supplied by the<br>manufacturer. A C&S report must<br>accompany this request.<br>*Demeclocycline will also be<br>approved for SIADH. |
| ULCERATIVE COLI            |   |  |   |
|                            |   | RAL  |   |
|                            | APRISO (mesalamine)<br>ASACOL (mesalamine) 400mg<br>COLAZAL (balsalazide)<br>DIPENTUM (olsalazine)<br>PENTASA (mesalamine) 250mg<br>sulfasalazine | ASACOL HD (mesalamine) 800mg<br>AZULFIDINE (sulfasalazine)<br>balsalazide<br>LIALDA (mesalamine)<br>PENTASA (mesalamine) 500mg   | Thirty (30) day trials of each of the<br>preferred agents of a dosage form<br>must be tried before a non-<br>preferred agent of that dosage form<br>will be authorized unless one of the<br>exceptions on the PA form is<br>present.  |



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|---------------------------|--|--|---|--|--|
|                           | RECTAL   |  |   |  |  |
|                           | CANASA (mesalamine)<br>mesalamine                                | SF ROWASA (mesalamine)   |   |  |  |
| VAGINAL ANTIBAC           | CTERIALS   |  |   |  |  |
|                           | clindamycin cream<br>METROGEL (metronidazole)                    | AVC (sulfanilamide)<br>CLEOCIN CREAM (clindamycin)<br>CLEOCIN OVULE (clindamycin)<br>CLINDESSE (clindamycin)<br>metronidazole<br>VANDAZOLE (metronidazole) | A trial, the duration of the<br>manufacturer's recommendation, of<br>each of the preferred agents is<br>required before a non-preferred<br>agent will be approved unless one<br>of the exceptions on the PA form is<br>present. |  |  |
| MISC BRAND/GEN            | ERIC   |  |   |  |  |
|                           | CLON   | IDINE  |   |  |  |
|                           | CATAPRES-TTS (clonidine)<br>clonidine tablets                    | clonidine patch<br>NEXICLON XR (clonidine)<br>CATAPRES TABLETS (clonidine)   | A thirty (30) day trial of each<br>preferred unique chemical entity in<br>the corresponding therapeutic<br>category is required before a non-<br>preferred agent will be authorized.  |  |  |
|                           |  | STROL  |   |  |  |
|                           | MEGACE ES (megestrol)<br>megestrol                               | MEGACE (megestrol)   |   |  |  |
|                           | SUBLINGUAL N   | IITROGLYCERIN  |   |  |  |
|                           | nitroglycerin sublingual<br>NITROSTAT SUBLINGUAL (nitroglycerin) | NITROLINGUAL (nitroglycerin)<br>NITROMIST (nitroglycerin)  |   |  |  |
|                           |  | EOTIDE   |   |  |  |
|                           | SANDOSTATIN (octreotide)   | octreotide   |   |  |  |
|                           | EPINEI   | PHRINE   |   |  |  |
|                           | TWINJECT (epinephrine)<br>EPIPEN (epinephrine)                   |  |   |  |  |
|                           |  | RACEPTIVES   |   |  |  |
|                           | YASMIN (ethinyl estradiol/drospirenone)                          | BEYAZ (ethinyl estradiol/drospirenone/levomefolate)  | 10  |  |  |



### PREFERRED DRUG LIST WITH PRIOR AUTHORIZATION CRITERIA

This is not an all-inclusive list of available covered drugs and includes only managed categories. Refer to cover page for complete list of rules governing this PDL.

04/01/12

**EFFECTIVE** 

| THERAPEUTIC<br>DRUG CLASS | PREFERRED AGENTS                            | NON-PREFERRED AGENTS   | PA CRITERIA  |
|---------------------------|---|--|--|
|                           |   | Gianvi (ethinyl estradiol/drospirenone)<br>Ocella (ethinyl estradiol/drospirenone)<br>YAZ (ethinyl estradiol/drospirenone) |  |
|                           | SUBSTANCE ABUSE TREATMENTS                  |  |  |
|                           | SUBOXONE (buprenorphine) FILM <sup>CL</sup> | SUBOXONE (buprenorphine) TABS  | Suboxone PA criteria is available at<br>http://www.dhhr.wv.gov/bms/Pharm<br>acy/Pages/pac.aspx |