Oral Tetracycline Derivatives
Prior Authorization
Program Summary

This program applies to FlexRx Open, FlexRx Closed, GenRx Open, GenRx Closed, Health Insurance Marketplace and KeyRx formularies.

FlexRx Standard and GenRx Standard prior authorization program.

The BCBS MN Step Therapy Supplement also applies to this program for all Commercial/HIM lines of business.

Program specific denial language for prerequisite step therapy component does not apply. Instead, supplemental program denial language will apply.

**FDA APPROVED INDICATIONS AND DOSAGE**

<table>
<thead>
<tr>
<th>Agent</th>
<th>Indications and Dosing</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Doxycycline Agents</strong></td>
<td></td>
</tr>
<tr>
<td>Acticlate™ (doxycycline hyclate tablet)</td>
<td>• For the treatment of infections due to susceptible strains of microorganisms (refer to labeling for additional details)</td>
</tr>
<tr>
<td>Adoxa®, Adoxa Pak® (doxycycline monohydrate tablet, capsule)</td>
<td>• In <em>acute intestinal amebiasis</em>, doxycycline may be a useful adjunct to amebicides</td>
</tr>
<tr>
<td>Doryx®, Doryx MPC® (doxycycline hyclate delayed-release tablet)</td>
<td>• For the prophylaxis of malaria due to <em>Plasmodium falciparum</em> in short-term travelers (&lt;4 months) to areas with chloroquine and/or pyrimethamine-sulfadoxine resistant strains</td>
</tr>
<tr>
<td>Doxycycline® (doxycycline hyclate delayed-release capsule; doxycycline hyclate tablet)</td>
<td>• In <em>severe acne</em>, doxycycline may be useful adjunctive therapy</td>
</tr>
<tr>
<td>Monodox®, Monodox® (doxycycline monohydrate capsule)</td>
<td>• Labeling does not include dosing recommendations for treatment of acne. Typical dosing is 50-100 mg twice daily</td>
</tr>
<tr>
<td>Targadox™ (doxycycline hyclate tablet)</td>
<td></td>
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<tr>
<td>Vibramycin® (doxycycline hyclate capsule)</td>
<td></td>
</tr>
<tr>
<td>Vibramycin® (doxycycline monohydrate suspension)</td>
<td></td>
</tr>
<tr>
<td>Vibramycin® (doxycycline calcium syrup)</td>
<td></td>
</tr>
<tr>
<td>Agent</td>
<td>Indications and Dosing</td>
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</tr>
<tr>
<td><strong>Oracea</strong> (doxycycline monohydrate delayed-release capsule)</td>
<td>• For the treatment of only inflammatory lesions (papules and pustules) of rosacea in adult patients. No meaningful effect was demonstrated for generalized erythema (redness) of rosacea. Oracea has not been evaluated for the treatment of the erythematous, telangiectatic, or ocular components of rosacea</td>
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- generic equivalents are available
- discontinued

<table>
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</tr>
<tr>
<td><strong>Minocin</strong> (minocycline capsule&lt;sup&gt;a&lt;/sup&gt;)</td>
<td>• For the treatment of infections due to susceptible strains of microorganisms (see labeling for details)</td>
</tr>
<tr>
<td><strong>minocycline tablet</strong></td>
<td>• In <em>acute intestinal amebiasis</em>, minocycline may be a useful adjunct to amebicides</td>
</tr>
<tr>
<td></td>
<td>• For the treatment of asymptomatic carriers of <em>Neisseria meningitidis</em> to eliminate meningococci from the nasopharynx</td>
</tr>
<tr>
<td></td>
<td>• In <em>severe acne</em>, minocycline may be useful adjunctive therapy</td>
</tr>
<tr>
<td></td>
<td>• Labeling does not include dosing recommendations for treatment of acne. Typical dosing is 50-100 mg twice daily.</td>
</tr>
</tbody>
</table>

<p>| <strong>Solodyn</strong>, <strong>Minocycline ER</strong> (minocycline extended-release tablet)&lt;sup&gt;a&lt;/sup&gt; | • To treat only inflammatory lesions of non-nodular moderate to severe acne vulgaris in patients 12 years of age and older. Solodyn did not demonstrate any effect on non-inflammatory lesions |
|                                                                              | • The recommended dosage of Solodyn is approximately 1 mg/kg once daily for 12 weeks |</p>
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</table>
| Minolira™ (minocycline extended-release tablet) | • To treat only inflammatory lesions of non-nodular moderate to severe acne vulgaris in patients 12 years of age and older. Minolira has not been evaluated in the treatment of infections  
  • The recommended dosage is approximately 1 mg/kg once daily for 12 weeks  
| Ximino™ (minocycline extended-release capsule)  | • To treat only inflammatory lesions of non-nodular moderate to severe acne vulgaris in patients 12 years of age and older  
  • Ximino did not demonstrate any effect on non-inflammatory acne lesions. Safety of Ximino has not been established beyond 12 weeks of use  
  • The recommended dosage of Ximino is approximately 1 mg/kg once daily for 12 weeks  |

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</table>
| Seysara™ (sarecycline tablets) | • To treat inflammatory lesions of non-nodular moderate to severe acne vulgaris in patients 9 years of age or older  
  • Limitations of Use: Efficacy of Seysara beyond 12 weeks and safety beyond 12 months have not been established. Seysara has not been evaluated in the treatment of infections. To reduce the development of drug-resistant bacteria as well as to maintain the effectiveness of other antibacterial drugs, Seysara should be used only as indicated.  
  • Dosage: The recommended dosage of is once daily with or without food.  
  60 mg for patients who weigh 33-54 kg  
  100 mg for patients who weigh 55-84 kg  
  150 mg for patients who weigh 85-136 kg  |

a – generic equivalents are available

**CLINICAL RATIONALE**  
**Acne Vulgaris**  
The American Academy of Dermatology suggests several options for treatment of acne vulgaris. Recommendations for topical acne therapies include benzoyl peroxide or combination of topical antibiotics (e.g. erythromycin or clindamycin) as monotherapy for mild acne, or in conjunction with topical retinoid, or systemic antibiotic therapy for moderate to severe acne. Topical antibiotics are not recommended in monotherapy due to
risk of bacterial resistance. Topical adapalene, tretinoin, and benzoyl peroxide can be safely used in the management of preadolescent acne in children. Azelaic acid is useful as an adjunctive acne treatment and is recommended in the treatment of postinflammatory dyspigmentation. Topical dapsone 5% gel is recommended for inflammatory acne, particularly in adult females with acne. There is limited data to support sulfur, nicotinamide, resorcinol, sodium sulfacetamide, aluminum chloride, and zinc in the treatment of acne.14

Systemic antibiotics have been a mainstay for acne treatment for years.14 They are indicated for use in moderate to severe inflammatory acne and should be used in combination with a topical retinoid and benzoyl peroxide. Tetracyclines are considered first-line therapy in moderate to severe acne, except when contraindicated. Doxycycline and minocycline are more effective than tetracycline but neither is superior to each other. Oral erythromycin and azithromycin should be reserved for those who cannot use tetracyclines. The use of other systemic antibiotics is discouraged due to limited data for use in acne. Trimethoprim-sulfamethoxazole and trimethoprim use should be restricted to patients who are unable to tolerate tetracycline or in treatment-resistant patients.14 Concomitant topical therapy with benzoyl peroxide or a retinoid should be used with systemic antibiotics and for maintenance after completion of systemic antibiotic therapy.14

Reviews of tetracycline agents used in the treatment of acne15,16 have found tetracycline, minocycline, and doxycycline all to be effective in the treatment of acne, particularly during the inflammatory stage. One review of seven randomized trials which were set up to compare the efficacy of tetracyclines found no evidence of superiority of one tetracycline over another in reducing acne lesion counts.15 Evidence-based recommendations for treatment of pediatric acne from the American Academy of Pediatrics consider oral antibiotics appropriate for moderate to severe inflammatory acne. Tetracycline derivatives, including tetracycline, doxycycline and minocycline are not to be used in children younger than 8 years of age.24

There are several other treatment options for acne. Hormonal therapy or oral contraceptives and isotretinoin are suggested; however, caution is needed for both therapies for adverse events and monitoring. There is limited evidence for the use and benefit of physical modalities for the routine treatment of acne, including pulsed dye laser, glycolic acid peels, and salicylic acid peels. Intraleisional corticosteroid injections are effective in the treatment of individual acne nodules. Furthermore, no current data supports any specific dietary changes to manage acne. However, data suggests that high glycemic index diets may be associated with acne and limited evidence suggests that some dairy products, particularly skim milk, may influence acne.14

**Rosacea**

Because there is no proven natural progression for rosacea, treatment choice is based on the patient’s current clinical manifestations and avoidance of known triggers is recommended.9

Topical agents are first-line therapy for the treatment of mild to moderate rosacea.21,22,23 Topical agents have less risk of adverse events, drug interactions and antibiotic resistance compared to systemic therapies.27 Topical metronidazole, azelaic acid, or brimonidine are recommended for erythema associated with rosacea. Topical ivermectin is recommended for rosacea inflammation with papulopustular lesions and may be used in combination with topical metronidazole or azelaic acid. Vascular laser therapy, such as pulsed dye laser, intense pulsed light, is also recommended for erythema and telangiectasia.9

Combination of topical antibiotics with oral antimicrobials could produce a more rapid response.23 Metronidazole and azelaic acid are standard topical antimicrobials used to treat the papules and pustules of rosacea; they appear to be about equally effective.21,23 Topical retinoids may be used for patients who do not respond to topical antimicrobials. Topical
Brimonidine is effective to treat moderate to severe erythema of rosacea. Systemic antibiotic therapy tends to be effective for treatment of papules, pustules, erythema and ocular inflammation. The severity of the patient's presentation helps guide the decision to initiate topical therapy alone or in combination with systemic therapy. Systemic therapy should be withdrawn when adequate response occurs.

Safety
The use of tetracycline agents, including doxycycline and minocycline, may cause fetal harm when administered during pregnancy. Drugs in the tetracycline class should not be used during pregnancy or by either gender when attempting to conceive. Photosensitivity reactions have been associated with tetracyclines. The use of drugs in the tetracycline class during tooth development may cause permanent tooth discoloration which is more common with long term use. Additionally, drugs in the tetracycline class have been associated with a reversible decrease in fibula growth rate caused by complexation with calcium. In general, tetracyclines should not be used in children under 8 years of age and extended-release minocycline should not be used in children under 12 years of age.

Minocycline
The safety and efficacy of Solodyn in the treatment of inflammatory lesions of non-nodular moderate to severe acne vulgaris was assessed in two 12-week, multi-center, randomized, double-blind, placebo-controlled, studies in subjects ≥12 years. The mean age of subjects was 20 years and subjects were from the following racial groups: white (73%), Hispanic (13%), black (11%), Asian/Pacific islander (2%), and other (2%). In the two efficacy and safety trials, a total of 924 subjects with non-nodular moderate to severe acne vulgaris received 1 mg/kg of Solodyn or placebo for a total of 12 weeks. The two primary efficacy endpoints were:

1. Mean percent change in inflammatory lesion counts from baseline to 12 weeks
2. Percentage of subjects with an Evaluator’s Global Severity Assessment (EGSA) of clear or almost clear at 12 weeks.

Patients on Solodyn had a greater mean percent improvement in inflammatory lesions (43.1% and 45.8% in studies one and two respectively) compared to placebo (31.7% and 30.8%) (p<0.05). Solodyn did not demonstrate any effect on non-inflammatory lesions.

There are no clinical studies comparing extended-release minocycline with older immediate-release formulations. A Medical Letter review of Solodyn concluded “Solodyn is an expensive new formulation of minocycline labeled for once-daily use. Whether Solodyn is as effective as immediate-release minocycline and less likely to cause vertigo remains to be established.”

Doxycycline
Oracea, indicated for the treatment of inflammatory lesions (papules and pustules) of rosacea in adult patients, is comprised of 30 mg immediate release and 10 mg delayed release doxycycline. While the mechanism of action is not fully understood, it is thought to be due to an anti-inflammatory effect.

The safety and efficacy of Oracea was evaluated in two double blind, randomized, placebo controlled trials involving 537 patients for the treatment of rosacea. Both phase III trials were 16 weeks in duration. Oracea therapy resulted in a mean decrease in lesion count from baseline of 11.8 and 9.5 in study one and two respectively compared to placebo (5.9 and 4.3) for placebo respectively (p<0.05). Patients on Oracea did not demonstrate improvement in erythema compared to placebo.

The FDA noted that the magnitude of efficacy shown is clinically somewhat limited and modest for an oral medication. The manufacturer has stated that at the systemic concentration provided by Oracea, doxycycline is not effective as an antimicrobial agent and...
appears to exert its action independent of antibacterial activity. The sponsor has not submitted data supporting this mechanism of action. Furthermore, there are some possible indicators of antibacterial action in the form of an increase in diarrhea in the active treatment arms of the pivotal trials.\textsuperscript{19}

A double-blind randomized trial compared Oracea 40 mg once daily to doxycycline 100 mg once daily in the treatment of moderate to severe rosacea for 16 weeks. There was no statistical significant difference in the primary efficacy endpoint of the change in total lesion count. There was a higher incidence of GI adverse events related to doxycycline 100 mg versus Oracea (26% vs 5%); however, the discontinuation rate was 50% higher with Oracea versus doxycycline 100 mg.\textsuperscript{20}

To receive an AB rating by the Food and Drug Administration (FDA) generic agents must be pharmaceutical equivalents to the innovator brand drug (contain the same active ingredients, are the same dosage form, the same route of administration, and are identical in strength or concentration) and the agent can be expected to have the same clinical effect and safety profile when administered to patients under the conditions specified in the innovator drug labeling.\textsuperscript{18} Generics may differ in shape, scoring, configuration, release mechanisms, packaging, excipients (including colors, flavors, preservatives), expiration time, and, within certain limits, labeling.\textsuperscript{10} AB-rated agents have had actual or potential bioequivalence problems resolved with adequate \textit{in vivo} and/or \textit{in vitro} evidence supporting bioequivalence.\textsuperscript{18} Doxycycline in oral capsules, oral tablets, and oral suspension and minocycline in oral capsules, oral tablets, and extended-release tablets are available as AB-rated generics.\textsuperscript{18}

For additional clinical information see Prime Therapeutics Formulary Chapter 1.4 Tetracyclines and Chapter 14.5 A-C Topical Acne Agents

REFERENCES
Oral Tetracycline Derivatives Prior Authorization

OBJECTIVE
The intent of the Oral Tetracycline Derivatives Prior Authorization (PA) criteria is to ensure appropriate selection of patients for treatment according to agent labeling and/or clinical studies and/or guidelines, and to encourage use of two first-line preferred oral agents – both doxycycline and minocycline – before use of these agents. Requests for nonpreferred agents will be reviewed when patient-specific documentation has been provided.

TARGET AGENTS

Doxycycline Agents:
- Acticlate™ (doxycycline hyclate tablet)
- Adoxa® (doxycycline monohydrate tablet, capsule)
- Doryx®, Doryx MPC® (doxycycline hyclate delayed-release tablet)*
- Doxycycline (doxycycline hyclate delayed-release capsule, doxycycline hyclate tablet, doxycycline monohydrate delayed release capsule)
- Monodox® (doxycycline monohydrate capsule)
- Oracea® (doxycycline monohydrate delayed-release capsule)
- Targadox™ (doxycycline hyclate tablet)
- Vibramycin® (doxycycline hyclate capsule, monohydrate suspension, doxycycline calcium syrup)

Minocycline Agents:
- Minocin® (minocycline capsule)
- minocycline tablet
- Minocycline ER (minocycline extended-release tablet)
- Minolira™ (minocycline extended-release tablet)
- Solodyn® (minocycline extended-release tablet)*
- Ximino™ (minocycline extended-release capsule)
  - a - available as a generic; generic designated target
  - b – discontinued

Tetracycline Agents:
- Seysara™ (sarecycline tablets)

PRIOR AUTHORIZATION CRITERIA FOR APPROVAL

Targeted agents will be approved when ALL of the following are met:

1. The patient has an FDA-labeled indication for the requested agent
   AND
2. The patient’s age is within FDA label for the requested indication
   AND
3. IF the patient’s diagnosis is acne or rosacea, ONE of the following:
   a. The patient is not currently being treated with another oral antibiotic for the treatment of acne or rosacea
   OR
   b. The patient is currently being treated with another oral antibiotic for the treatment of acne or rosacea and the antibiotic will be discontinued before starting the requested agent
   AND
4. The patient’s medication history includes use of a preferred oral generic doxycycline agent in the past 180 days OR the patient has a documented intolerance, FDA labeled contraindication, or hypersensitivity to a preferred oral generic doxycycline agent
   AND
5. The patient’s medication history includes use of a preferred oral generic minocycline agent in the past 180 days OR the patient has a documented intolerance, FDA labeled contraindication, or hypersensitivity to a preferred oral generic minocycline agent

**Length of Approval:** 12 months
Step Therapy Supplement

This program applies to FlexRx Closed, FlexRx Open, GenRx Closed, GenRx Open, Health Insurance Marketplace, FocusRx and KeyRx formularies.

Please note, this does not include or apply to quantity limit questions.

STEP THERAPY SUPPLEMENT

OBJECTIVE
The intent of the Step Therapy Supplement is to provide additional questions, to ensure compliance to MN Statute 62Q.184. These questions will apply if the step therapy component within a Prior Authorization or Step Therapy program is not able to be approved.

CONDITIONS FOR APPROVAL
The requested agent will be approved when ONE of the following are met:

1. The patient is currently being treated with the requested agent as indicated by ALL of the following:
   a. A statement by the prescriber that the patient is currently taking the requested agent
   AND
   b. A statement by the prescriber that the patient is currently receiving a positive therapeutic outcome on requested agent
   AND
   c. The prescriber states that a change in therapy is expected to be ineffective or cause harm

OR

2. The patient’s medication history include the required prerequisite/preferred agent(s) as indicated by:
   a. Evidence of a paid claim(s) within the past 999 days
   OR
   b. The prescriber has stated that the patient has tried the required prerequisite/preferred agent(s) in the past 999 days AND the required prerequisite/preferred agent(s) was discontinued due to lack of effectiveness or an adverse event

OR

3. The prescriber has provided documentation that the required prerequisite/preferred agent(s) cannot be used due to a documented medical condition or comorbid condition that is likely to cause an adverse reaction, decrease ability of the patient to achieve or maintain reasonable functional ability in performing daily activities or cause physical or mental harm

Length of Approval: As per program specific criteria