Louisiana Medicaid Hemodialysis – Phosphate Binders

The *Louisiana Uniform Prescription Drug Prior Authorization Form* should be utilized to request prior authorization for non-preferred phosphate binders.

Additional Point-of-Sale edits may apply.

By submitting the authorization request, the prescriber attests to the conditions available HERE.

These agents may have Black Box Warnings and/or may be subject to Risk Evaluation and Mitigation

Strategy (REMS) under FDA safety regulations. Please refer to individual prescribing information for details.

Approval Criteria for Initiation and Continuation of Therapy

Approval Criteria for Initial and Reauthorization Requests

- For sevelamer tablets (generic for Renvela®) there has been a treatment failure or intolerable side effect with or contraindication to brand Renvela®; **OR**
- There is no preferred alternative that is the exact same chemical entity, formulation, strength, and delivery device; **AND**
- Previous use of a preferred product **ONE** of the following is required:
 - The recipient has had a treatment failure with at least one preferred product; **OR**
 - The recipient has had an *intolerable side effect* to at least one preferred product; **OR**
 - \circ The recipient has *documented contraindication(s)* to all of the preferred products that are appropriate to use for the condition being treated; **OR**
 - \circ There is *no preferred product that is appropriate* to use for the condition being treated; **OR**
 - The prescriber states that the recipient is currently using the requested medication AND one of the following applies:
 - There is evidence in pharmacy claims of at least 60 days of the requested medication within the previous 90-day period; OR
 - There is evidence in pharmacy claims of less than 60 days of the requested medication AND the prescriber states the recipient has been treated with the requested medication in an inpatient facility; OR
 - There is evidence in pharmacy claims of less than 60 days of the requested medication **AND** the prescriber has verified that the pharmacy has dispensed at least 60 days of medication (billed to other insurance, and therefore not viewable in pharmacy claims).; **AND**
- By submitting the authorization request, the prescriber attests to the following:
 - The prescribing information for the requested medication has been thoroughly reviewed, including any Black Box Warning, Risk Evaluation and Mitigation Strategy (REMS), contraindications, minimum age requirements, recommended dosing, and prior treatment requirements; AND
 - All laboratory testing and clinical monitoring recommended in the prescribing information have been completed as of the date of the request and will be repeated as recommended; AND-
 - The recipient has no concomitant drug therapies or disease states that limit the use of the requested medication and will not be receiving the requested medication in

combination with any other medication that is contraindicated or not recommended per FDA labeling.

<u>Duration of approval for initiation and continuation of therapy</u>Duration of initial and reauthorization approval: 12 months

References

Clinical Pharmacology [database online]. Tampa, FL: Gold Standard, Inc.; Retrieved from https://www.clinicalkey.com/pharmacology/

DiPiro JT, Talbert RL, Yee GC, Matzke GR, Wells BG, Posey L. eds. Pharmacotherapy: A Pathophysiologic Approach, 10e New York, NY: McGraw-Hill; Retrieved from https://accesspharmacy.mhmedical.com/book.aspx?bookid=1861

Revision / Date	Implementation Date
Single PDL Implementation	May 2019
Added specific wording for use of RenaGel®, separated "Select	
Therapeutic Classes with Established Recent Claims" into	January 2020
individual therapeutic class documents / November 2019	-
Removed wording requiring use of brand RenaGel® and removed	July 2020
reference, formatting changes / July 2020	July 2020
Added specific wording for use of Renvela® tablets, formatting	Index 2021
changes / April 2021	July 2021
Removed specific wording for the use of Renvela® tablets,	July 2024
formatting changes / April 2024	<u>July 2024</u>