Template Letter of Appeal

This Template Letter of Appeal is designed to assist with your payer interactions concerning coverage or reimbursement for RUCONEST (C1 esterase inhibitor [recombinant]) in the event of a coverage or claim denial. Use of this document does not guarantee coverage or reimbursement. This letter is designed as a framework only, and should be customized appropriately by your facility.

As a healthcare professional, you are solely responsible for providing accurate information to third-party payers. The information in this document should accurately reflect your patient’s history and your clinical rationale as to why RUCONEST is medically necessary.

Indications for RUCONEST

RUCONEST® is a C1 esterase inhibitor (recombinant) indicated for the treatment of acute attacks in adult and adolescent patients with hereditary angioedema (HAE). Effectiveness in clinical studies was not established in HAE patients with laryngeal attacks.

Important Safety Information for RUCONEST

RUCONEST® (C1 esterase inhibitor [recombinant]) is contraindicated in patients with a history of allergy to rabbits or rabbit-derived products, and patients with a history of life-threatening immediate hypersensitivity reactions to C1 esterase inhibitor preparations, including anaphylaxis.

Severe hypersensitivity reactions may occur. The signs and symptoms of hypersensitivity reactions may include hives, generalized urticaria, tightness of the chest, wheezing, hypotension, and/or anaphylaxis during or after injection of RUCONEST. Should symptoms occur, discontinue RUCONEST and administer appropriate treatment. Because hypersensitivity reactions may have symptoms similar to HAE attacks, treatment methods should be carefully considered.

Serious arterial and venous thromboembolic (TE) events have been reported at the recommended dose of plasma-derived C1 esterase inhibitor products in patients with risk factors. Risk factors may include the presence of an indwelling venous catheter/access device, prior history of thrombosis, underlying atherosclerosis, use of oral contraceptives or certain androgens, morbid obesity, and immobility. Monitor patients with known risk factors for TE events during and after RUCONEST administration.

RUCONEST has not been studied in pregnant women; therefore, should only be used during pregnancy if clearly needed.

The most common adverse reactions (incidence ≥2%) were headache, nausea, and diarrhea. The serious adverse reaction in clinical studies of RUCONEST was anaphylaxis.

Please see accompanying full Prescribing Information for RUCONEST or visit www.ruconest.com.
[Date]

[Insurance Company]
[Insurance Address]
[Policy Number]
[Group Number]
[Member ID]

Re: [Patient Name] [Case Reference ID]

Dear [Contact/Department],

This is a formal letter of appeal for reconsideration of coverage for RUCONEST for [Patient Name]. [Patient Name] has been diagnosed and has suffered from hereditary angioedema (HAE) for [insert # months/years]. [Payer Name] has stated that RUCONEST is not covered because [Denial Reason]. As the treating physician, it is my clinical judgment that RUCONEST is the most medically appropriate treatment for [Patient Name’s] acute HAE attacks. It is inappropriate for [Payer Name] to deny coverage of RUCONEST given my patient has had continued [breakthrough attacks on their existing therapy(ies), needs a C1-INH that returns their C1 level to a normal range quickly, etc.] [Patient Name] is still at risk for continued attacks, hospitalizations, and suffering from this disease.

Patient History and Diagnosis
[Patient Name] is a [age] year-old [male/female] and has been hospitalized [# of times] due to [his/her] frequent HAE attacks. This has resulted in debilitating HAE attacks, which require immediate medical intervention. My patient has an average of [#] HAE attacks per [week/month/year]. These HAE attacks are difficult to manage due to the inability to predict the location or severity of the attack.

Given [Patient’s Name] clinical history and documented treatments tried and failed, RUCONEST should be covered for this patient’s condition. [My patient has had the opportunity to try RUCONEST, and has responded positively. I believe this is the most appropriate acute therapy for my patient to use in conjunction with X drug(s).]

To support this appeal for coverage, I have included the following documentation:

- [Denial letter or payer communication]
- [Initial LMN/PA form]
- [Treatment history, therapies tried and failed, reason for failure, date of discontinuation and/or reason why existing therapy is not enough, length of time]
- [Frequency of attacks, date of last attack, level of severity]
- [Describe areas predominantly affected (abdomen, extremities, face, urogenital tract, other)]
- [Details from patient diary documenting patient attacks]
- [Positive response to RUCONEST trial units]
- [Images provided by patients during an HAE attack]
- [HAEA Practice Guidelines]
- [Relevant clinical reprints]
- [RUCONEST Prescribing Information]

Additional Information regarding RUCONEST
• RUCONEST is a plasma-free, recombinant product that is indicated for the treatment of acute attacks in adult and adolescent patients diagnosed with hereditary angioedema (HAE). Efficacy has not been established in laryngeal attacks.  

• As a recombinant product, RUCONEST carries no risk of blood-borne infections or potential drug shortages.

• RUCONEST is an intravenous (IV) injectable which may be self-administered by appropriately trained patients at a dose of 50 IU/kg with a maximum of 4200 IU per dose, to be given as a slow IV injection.

• RUCONEST has demonstrated rapid symptom relief, as well as sustained response for at least 3 days in most patients; thus, the patient may not require repeat therapy during this time frame.

• RUCONEST directly addresses the cause of HAE by replacing and/or replenishing the inadequate C1-INH levels produced by the patient's body.

• The United States HAE Association guidelines recommend that an effective treatment strategy for acute HAE includes replenishing C1-INH levels as soon as possible at the onset of an attack.

• RUCONEST is contraindicated in patients with a history of allergy to rabbits or rabbit-derived products, and patients with a history of life-threatening immediate hypersensitivity reactions to C1 esterase inhibitor preparations, including anaphylaxis.

• Please see accompanying full Prescribing Information for RUCONEST or visit www.ruconest.com.

On behalf of [Patient Name], we respectfully request you overturn your negative coverage decision for RUCONEST. Please call me at [Primary Treating Site Phone Number] if I can be of further assistance or if you require additional information.

Sincerely,

[treating provider name]