

PHARMACY POLICY – 5.01.584


CGRP Inhibitors for Migraine Prophylaxis

Effective Date: Jan. 1, 2019
Last Revised: Dec. 13, 2018
Replaces: N/A

RELATED PHARMACY / MEDICAL POLICIES:
5.01.503 Migraine and Cluster Headache Medications

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Introduction

Migraine is a debilitating disease, with severe headaches. Some people have other symptoms like seeing auras, experiencing nausea or vomiting, and suffering an inability to tolerate bright light or loud noises. About one in eight Americans has migraines. It's the seventh most disabling disease worldwide. Women are twice as likely as men to suffer from migraine.

Some people have just a few headaches a month. These may be treated with pills like ibuprofen or prescription medications. These treatments stop the headaches after they've started. However, if people take too much of the headache-stopping medications, over time they may end up with more headaches. This is a poor long-term strategy. It's estimated that more than 40% of migraine patients have unmet needs. These include experiencing disability during a migraine attack, not being happy with existing treatments, or overusing habit-forming medications. Newer types of migraine-preventing drugs have been developed to address these unmet needs. This policy describes when these types of drugs may be considered medically necessary.

Note: The Introduction section is for your general knowledge and is not to be taken as policy coverage criteria. The rest of the policy uses specific words and concepts familiar to medical professionals. It is intended for providers. A provider can be a person, such as a doctor, nurse, psychologist, or dentist. A provider also can be a place where medical care is given, like a hospital, clinic, or lab. This policy informs them about when a service may be covered.

Policy Coverage Criteria

Drug	Medical Necessity
Aimovig™ (erenumab) Ajovy™ (fremanezumab) Emgality™ (galcanezumab)	Aimovig™ (erenumab), Ajovy™ (fremanezumab), or Emgality™ (galcanezumab) may be considered medically necessary in patients who have: <ul style="list-style-type: none"> An average of 5 or more migraine days per month AND <ul style="list-style-type: none"> Tried three different categories of prophylactic migraine headache therapies listed in the Appendix section AND <ul style="list-style-type: none"> Tried three different triptan medications for abortive use unless triptan medications are contraindicated

Length of Approval	
Approval	Criteria
Initial Approval	Initial approval will be for three months.
Reauthorization	Continued therapy will be approved for periods of up to one year as long as the patient has shown and continues to show a sustained reduction in headache frequency OR severity compared to baseline prior to initiation of treatment with the CGRP inhibitor.
Required Documentation	Chart notes describing patient’s diagnosis and progress including headache frequency; medication history, if not documented by our prescription claims record.

Coding

N/A

Related Information



Benefit Application

All CGRP inhibitors are managed through the pharmacy benefits.

Evidence Review

Description

Medical Condition

Migraine is a debilitating neurologic disorder, characterized by severe headache that can be accompanied by aura and additional symptoms such as nausea, vomiting, photosensitivity, phonophobia, visual disturbance, and tingling in the hands and feet. Migraine affects 12-16% of the US population and is the seventh most disabling disease worldwide. It disproportionately affects females, with females 2-2.5 times more likely to suffer from migraine.

The frequency of migraine attacks a patient experiences guides migraine treatment approach. Low frequency of attacks (1-3 migraines per month) can be treated with abortive therapy, such as triptans, NSAIDs, and other analgesics. However, sufferers of episodic migraine, with 4-14 migraine days per month, and chronic migraine, with >15 headache days per month (8 of which are migraine), should receive preventative migraine therapy. Individual patient response to migraine therapy is highly variable, with over 23 genes purported to be linked to the neuronal and vascular pathophysiology of migraine, and effective migraine treatment options to date are limited.

As few therapies are effective in preventing migraine, drug tolerance and subsequent overuse of abortive therapies is common and carry the risk of causing rebound headaches. Thus, these are not a viable long-term treatment option. Over 40% of patients with migraines have unmet needs, including disability, treatment dissatisfaction, and opioid/barbiturate overuse or dependence. Thus, new treatments, both for prophylaxis and acute therapy, are eagerly awaited to help alleviate the burden of migraine-related disability.

Migraine has a significant economic and social impact in the U.S. During a migraine attack, sufferers of migraine are often bedridden and have severe impairment to their ability to function.



Migraineurs frequently isolate themselves in a quiet, dark area and can lose several days of productivity while experiencing substantial pain. This, in turn, can lead to further comorbid conditions—most commonly depression—which contribute to the economic burden as well. The economic impact on both employers due to loss of productivity and the healthcare systems due to utilization, are estimated to be over 13 billion dollars.

Treatment Alternatives

Migraine treatment consists of two pillars: abortive treatment for acute migraine attacks and prophylactic treatment. While there are many numerous medications indicated for use in chronic and episodic migraine prophylaxis, there are currently no reliable treatments for the prevention of migraine, as all are drugs designed for targets other than migraine. Medications approved for the prevention of chronic and episodic migraine include beta-blockers, antiepileptic drugs, and antidepressants. The most commonly used medications are propranolol, topiramate and amitriptyline. Due to low efficacy in migraine prevention and the side effect profile, adherence is poor, with about 26% of patients at 6 months and 17% of patients at 12 months continuing these drugs.

Triptans, via inhibition of 5-hydroxytryptamine receptors in the intracranial vasculature, are effective for treating acute migraine, and can be used to prevent progression if administered in the early stage of migraine. However, tolerance to these medications occurs quickly and overuse contributes to headache exacerbation. Triptan quantity limits are typically set to accommodate treatment of three migraine days per month. NSAIDs and ergotamines can also be used as abortive therapies, though as with triptans, rebound headaches from overuse can occur with these as well.

Based on the data published from the PREEMPT trials I and II in 2011, onabotulinumtoxinA (Botox) was approved for use in chronic migraine despite numerous studies conducted previously failed to show a statistically significant difference between Botox™ and placebo in reduction of migraine days. Since, incobotulinumtoxinA (Xeomin™) and abobotulinumtoxinA (Dysport™) have been approved for the treatment of CM, however, failure to respond and development of antibodies to the botulinum toxin, in addition to the specialized administration these injections require, prevent these drugs from being a viable migraine prevention treatment for many patients.



Aimovig™ (erenumab)

Aimovig™ (erenumab) is a fully human monoclonal antibody that targets the CGRP receptor through competitive, reversible inhibition. Erenumab binds the CGRP receptor with high specificity and potency, blocking the action of CGRP and prevent vasodilation.

Ajovy™ (fremanezumab)

Fremanezumab is a humanized monoclonal antibody that targets the CGRP receptor through competitive, reversible inhibition. Like Erenumab, fremamanezumab binds the CGRP receptor with high specificity and potency, blocking the action of CGRP and prevent vasodilation.

Emgality™ (galcanezumab)

Like fremanezumab, galcanezumab is a humanized monoclonal antibody that binds to the CGRP receptor. Erenumab, fremanezumab and galcanezumab all bind the CGRP receptor with high specificity and potency, blocking the action of CGRP and prevent vasodilation.

Rationale

Efficacy/Effectiveness

Aimovig™ (erenumab) is a fully human monoclonal antibody that targets the calcitonin-gene-related peptide receptor to prevent both episodic and chronic migraine. Evidence for efficacy of erenumab in the prevention of episodic migraine was evaluated in STRIVE, a Phase III, randomized controlled trial (RCT), which included 955 patients with history of episodic migraine; STRIVE observed a reduction of 3.2 and 3.7 migraine days per month for patients treated with erenumab 70 mg and erenumab 140 mg, respectively, compared to a 1.8 migraine day per month reduction in the placebo group.

Evidence for efficacy of erenumab in the treatment of chronic migraine was evaluated in a Phase II randomized controlled trial (RCT), which included 656 patients with history of chronic migraine and observed a mean reduction in mean monthly migraine days (MMMD) of 6.6 days in patients treated with erenumab 70mg, 6.6 days in patients treated with erenumab 140mg, and 4.2 days in placebo group.



Though no studies yet show the long-term efficacy and durability of treatment with CGRP monoclonal antibodies beyond one year, efficacy has been demonstrated in patients treated with erenumab 70 mg and erenumab 140 mg every 4 weeks for 6 months. Additionally, an ongoing open-label extension has evaluated the efficacy of erenumab 70 mg every 4 weeks at one year.

In STRIVE, 50% of patients receiving erenumab 140 mg and 43.3% of patients receiving erenumab 70 mg were able to achieve >50% reduction in MMMD. In the OLE, patients were able to reduce MMMD from 8.8 days at baseline, to 6.3 days at the end of the 12-week double-blind period, to 3.7 days at the one-year interim analysis. Due to this level of efficacy at one year, erenumab is likely to significantly improve the quality of life for patients within the episodic migraine population.

Ajovy™ (fremanezumab) is a humanized IgG2a monoclonal antibody that selectively binds both α and β isoforms of the calcitonin gene-related peptide (CGRP). Evidence for efficacy of fremanezumab in the prevention of chronic migraine was evaluated in HALO CM, a phase III, randomized controlled trial (RCT), which included 1130 patients with chronic migraine. HALO CM observed a reduction of 4.9 and 5.0 headache days per month for patients treated with fremanezumab 675 mg quarterly and fremanezumab 675 mg followed by a monthly fremanezumab dose of 225 mg, respectively, compared to a reduction of 3.2 headache days per month in the placebo group. Thirty-eight percent of patients in the fremanezumab quarterly group and 41% of patients in the fremanezumab monthly group attained at least 50% reduction in the average number of headache days per month, compared to 18% of patients in the placebo group.

Evidence for efficacy of fremanezumab in the treatment of high-frequency episodic migraine was evaluated in a phase II randomized controlled trial (RCT), which included 297 patients with at most 14 headache days but at least 8 migraine days per month. The observed mean reduction in mean monthly migraine days (MMMD) from baseline to weeks 9 – 12 was 6.27 days in patients treated with fremanezumab 225mg monthly and 6.09 days in patients treated with fremanezumab 675 mg monthly, compared to 3.46 days in placebo group.

Further evidence for efficacy of fremanezumab in the treatment of episodic migraine is being evaluated the phase III HALO EM trial but the results have not been published in any peer-reviewed journal.

Galcanezumab was studied in the EVOLVE-1 and EVOLVE-2 Phase 3, double-blind, randomized-controlled trials that compared galcanezumab at 120 mg and 240 mg to placebo for six months in patients diagnosed with episodic migraine. The main difference between the two trials was that EVOLVE-1 observed patients in North America only, while EVOLVE-2



observed patients internationally. In EVOLVE-1, patients saw significant benefits in the galcanezumab 120 mg and 240 mg groups with a reduction of 4.7 and 4.6 monthly migraine headache days (MHDs), respectively, compared to placebo with a reduction of 2.8 monthly MHDs ($p < 0.001$ for both comparisons). Similar improvements were seen in EVOLVE-2; the galcanezumab 120 mg and 240 mg groups saw a reduction of 4.3 and 4.2 monthly MHDs, respectively, compared to placebo with a reduction of 2.3 monthly MHDs ($p < 0.001$ for both comparisons). For EVOLVE-1 and EVOLVE-2, galcanezumab further demonstrated improvements in terms of monthly MHDs with acute migraine medication use, Migraine-Specific Quality of Life Role-Functioning Restrictive (MSQ RF-R) scores, and percent reductions in monthly MHDs (>50%, >75%, and >100%).

REGAIN is an unpublished Phase 3, double-blind, randomized-controlled trial that compared galcanezumab at 120 mg and 240 mg to placebo for three months in patients diagnosed with chronic migraine. Both galcanezumab groups saw significant improvement with reductions of 4.83 and 4.62 monthly MHDs, respectively, compared to placebo with a reduction of 2.47 monthly MHD ($p < 0.001$ for both comparisons). 50% and 75% reductions in monthly MHDs had significant improvements in both galcanezumab groups. <2% of patients saw a 100% reduction in monthly MHDs with no significant differences between groups. Changes in the number of monthly MHDs with acute migraine medication use and MSQ RF-R scores showed improvements in both galcanezumab groups.

REBUILD is an ongoing trial studying galcanezumab in ages 6-17, and CONQUER is another ongoing trial studying galcanezumab in patients who have failed previous migraine preventive medications in the past 10 years due to inadequate efficacy or tolerability. A long-term open-label study is also being conducted in patients with episodic or chronic migraines. Once available, these studies may expand galcanezumab's applicability and generalizability.

Safety/Tolerability

The overall safety profile of erenumab was similar to that of placebo in clinical trials. The 28 week open-label continuation study with STRIVE patients is still underway, results from which will be required to assess long term safety of erenumab. In the interim analysis of the OLE from phase II trial, 13.1% of patients developed drug-binding antibodies, and 2.4% developed drug-neutralizing antibodies; this led to a transient response in a total of 37 patients.

A larger portion of patients treated with fremanezumab had a treatment-related adverse event than those in the placebo group. However, the most common adverse events were injection-site reactions (pain, induration, and erythema), which should not have any significant impact on the



safety of the product. There is no published study evaluating long-term efficacy and safety of fremanezumab. The ongoing phase III FOCUS trial will evaluate the efficacy and safety as a migraine prophylaxis up to 24 weeks (with the last 12 weeks as open label). Another ongoing phase III trial in patients with cluster headaches may shed light on adverse events with longer use up to 36 weeks.

The overall safety profile of galcanezumab was slightly worse than that of placebo in clinical trials. In EVOLVE-1, the rate of patients reporting >1 treatment-emergent adverse event (TEAE) was reported for 65.5% and 67.7% of patients in the 120 mg and 240 mg groups, compared to 60.4% in the placebo group. EVOLVE-2 saw 65% and 71.5% of patients in the 120 mg and 240 mg groups, respectively, report 1 > TEAE, compared to 62.3% of patients in the placebo group. Serious adverse events (SAEs) were rare and uncommon in galcanezumab. In some cases, investigators determined that SAEs were not associated with the drug. Injection-site related adverse events were the most frequent adverse event (AE) in all the trials, ranging from 6-20%, but none were significantly different from the placebo group. Neutralizing anti-drug antibodies (ADAs) were reported in 18/415 patients in the EVOLVE-1 trial and 29 patients in EVOLVE-2 (EVOLVE-2's rate included both placebo and galcanezumab). Galcanezumab was well-tolerated with about 1-4% of patients discontinuing due to adverse events, but investigators did not specify what these adverse events were.

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33. Policy was reviewed by a board certified practicing neurologist with specialty in headache management. Approved by the independent P&T Committee May 30, 2018.

Appendix

Prophylactic Therapy

Antidepressants, Beta blockers, Calcium channel blockers, Naproxen, Ergotamine preparations, Divalproex sodium, Topiramate, Botulinum toxin (Botox®), Calcitonin gene-related peptide (CGRP) antagonists, Others (cyproheptadine, clonidine, other anticonvulsants)

History

Date	Comments
06/01/18	New policy, approved May 17, 2018. Add to Prescription Drug section. Aimovig™ (erenumab) may be considered medically necessary in patients with an average of more than 4 migraine days per month who have failed at least three preventive migraine therapies and are receiving a maximum monthly supply of abortive migraine treatments.



Date	Comments
10/01/18	Interim Review, approved September 20, 2018. Added criteria for Ajovy (fremanezumab).
01/01/19	Interim Review, approved December 13, 2018. Added criteria for Emgality (galcanezumab).

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本通知有重要的訊息。本通知可能有關於您透過 Premera Blue Cross 提交的申請或保險的重要訊息。本通知內可能有重要日期。您可能需要在截止日期之前採取行動，以保留您的健康保險或者費用補貼。您有權利免費以您的母語得到本訊息和幫助。請撥電話 800-722-1471 (TTY: 800-842-5357)。

Oromoo (Cushite):

Beeksisni kun odeeffannoo barbaachisaa qaba. Beeksisti kun sagantaa yookan karaa Premera Blue Cross tiin tajaajila keessan ilaalchisee odeeffannoo barbaachisaa qabaachuu danda'a. Guyyaawwan murteessaa ta'an beeksisa kana keessatti ilaalaa. Tarii kaffaltiidhaan deeggaramuuf yookan tajaajila fayyaa keessaniif guyyaa dhumaa irratti wanti raawwattan jiraachuu danda'a. Kaffaltii irraa bilisa haala ta'een afaan keessaniin odeeffannoo argachuu fi deeggarsa argachuuf mirga ni qabaattu. Lakkoofsa bilbilaa 800-722-1471 (TTY: 800-842-5357) tii bilbilaa.

Français (French):

Cet avis a d'importantes informations. Cet avis peut avoir d'importantes informations sur votre demande ou la couverture par l'intermédiaire de Premera Blue Cross. Le présent avis peut contenir des dates clés. Vous devez peut-être prendre des mesures par certains délais pour maintenir votre couverture de santé ou d'aide avec les coûts. Vous avez le droit d'obtenir cette information et de l'aide dans votre langue à aucun coût. Appelez le 800-722-1471 (TTY: 800-842-5357).

Kreyòl ayisyen (Creole):

Avi sila a gen Enfòmasyon Enpòtan ladann. Avi sila a kapab genyen enfòmasyon enpòtan konsènan aplikasyon w lan oswa konsènan kouvèti asirans lan atravè Premera Blue Cross. Kapab genyen dat ki enpòtan nan avi sila a. Ou ka gen pou pran kèk aksyon avan sèten dat limit pou ka kenbe kouvèti asirans sante w la oswa pou yo ka ede w avèk depans yo. Se dwa w pou resewva enfòmasyon sa a ak asistans nan lang ou pale a, san ou pa gen pou peye pou sa. Rele nan 800-722-1471 (TTY: 800-842-5357).

Deutsche (German):

Diese Benachrichtigung enthält wichtige Informationen. Diese Benachrichtigung enthält unter Umständen wichtige Informationen bezüglich Ihres Antrags auf Krankenversicherungsschutz durch Premera Blue Cross. Suchen Sie nach eventuellen wichtigen Terminen in dieser Benachrichtigung. Sie könnten bis zu bestimmten Stichtagen handeln müssen, um Ihren Krankenversicherungsschutz oder Hilfe mit den Kosten zu behalten. Sie haben das Recht, kostenlose Hilfe und Informationen in Ihrer Sprache zu erhalten. Rufen Sie an unter 800-722-1471 (TTY: 800-842-5357).

Hmoob (Hmong):

Tsawb ntawv tshaj xo no muaj cov ntshiab lus tseem ceeb. Tej zaum tsawb ntawv tshaj xo no muaj cov ntshiab lus tseem ceeb txog koj daim ntawv thov kev pab los yog koj qhov kev pab cuam hnu ntawm Premera Blue Cross. Tej zaum muaj cov hnuv tseem ceeb uas sau rau hauv daim ntawv no. Tej zaum koj kuj yuav tau ua qee yam uas peb kom koj ua tsis pub dhau cov caij nyoog uas teev tseg rau hauv daim ntawv no mas koj thiaj yuav tau txais kev pab cuam kho mob los yog kev pab them tej nqi kho mob ntawd. Koj muaj cai kom lawv muab cov ntshiab lus no uas tau muab sau ua koj hom lus pub dawb rau koj. Hu rau 800-722-1471 (TTY: 800-842-5357).

Iloko (Ilocano):

Daytoy a Pakdaar ket naglaon iti Napateg nga Impormasion. Daytoy a pakdaar mabalin nga adda ket naglaon iti napateg nga impormasion maipanggep iti aplikasyonyo wenna coverage babaen iti Premera Blue Cross. Daytoy ket mabalin dagiti importante a petsa iti daytoy a pakdaar. Mabalin nga adda rumbeng nga aramidenyo nga addang sakbay dagiti partikular a naituding nga aldaw tapno mapagtalinaedyo ti coverage ti salun-ato wenna tulong kadagiti gastos. Adda karbenganyo a mangala iti daytoy nga impormasion ken tulong iti bukodyo a pagsasao nga awan ti bayadanyo. Tumawag iti numero nga 800-722-1471 (TTY: 800-842-5357).

Italiano (Italian):

Questo avviso contiene informazioni importanti. Questo avviso può contenere informazioni importanti sulla tua domanda o copertura attraverso Premera Blue Cross. Potrebbero esserci date chiave in questo avviso. Potrebbe essere necessario un tuo intervento entro una scadenza determinata per consentirti di mantenere la tua copertura o sovvenzione. Hai il diritto di ottenere queste informazioni e assistenza nella tua lingua gratuitamente. Chiama 800-722-1471 (TTY: 800-842-5357).

日本語 (Japanese):

この通知には重要な情報が含まれています。この通知には、Premera Blue Cross の申請または補償範囲に関する重要な情報が含まれている場合があります。この通知に記載されている可能性がある重要な日付をご確認ください。健康保険や有料サポートを維持するには、特定の期日までに行動を取らなければならない場合があります。ご希望の言語による情報とサポートが無料で提供されます。800-722-1471 (TTY: 800-842-5357)までお電話ください。

한국어 (Korean):

본 통지서에는 중요한 정보가 들어 있습니다. 즉 이 통지서는 귀하의 신청에 관하여 그리고 Premera Blue Cross 를 통한 커버리지에 관한 정보를 포함하고 있을 수 있습니다. 본 통지서에는 핵심이 되는 날짜들이 있을 수 있습니다. 귀하의 건강 커버리지를 계속 유지하거나 비용을 절감하기 위해서 일정한 마감일까지 조치를 취해야 할 필요가 있을 수 있습니다. 귀하의 이러한 정보와 도움을 귀하의 언어로 비용 부담없이 얻을 수 있는 권리가 있습니다. 800-722-1471 (TTY: 800-842-5357) 로 전화하십시오.

ລາວ (Lao):

ແຈ້ງການນີ້ມີຂໍ້ມູນສໍາຄັນ. ແຈ້ງການນີ້ອາດຈະມີຂໍ້ມູນສໍາຄັນກ່ຽວກັບຄໍາຮ້ອງສະໝັກ ຫຼື ຄວາມຄົມຄອງປະກັນໄພຂອງທ່ານຜ່ານ Premera Blue Cross. ອາດຈະມີວັນທີ່ສໍາຄັນໃນແຈ້ງການນີ້. ທ່ານອາດຈະຈຳເປັນຕ້ອງດໍາເນີນການຕາມກຳນົດ ເວລາສະເພາະເພື່ອຮັກສາຄວາມຄົມຄອງປະກັນສະພາບ ຫຼື ຄວາມຊ່ວຍເຫຼືອເວັ້ນເວົ້ອງຄ່າໃຊ້ຈ່າຍຂອງທ່ານໄດ້. ທ່ານມີສິດໄດ້ຮັບຂໍ້ມູນນີ້ ແລະ ຄວາມຊ່ວຍເຫຼືອເປັນພາສາຂອງທ່ານໂດຍບໍ່ເສຍຄ່າ. ໃຫ້ໃບທາ 800-722-1471 (TTY: 800-842-5357).

ភាសាខ្មែរ (Khmer):

សេចក្តីជូនដំណឹងនេះមានព័ត៌មានយ៉ាងសំខាន់។ សេចក្តីជូនដំណឹងនេះប្រហែលជាមានព័ត៌មានយ៉ាងសំខាន់អំពីទម្រង់បែបបទ ឬការរៀបចំរបស់អ្នកកាមរយ: Premera Blue Cross ។ ប្រហែលជាមាន កាលបរិច្ឆេទសំខាន់នៅក្នុងសេចក្តីជូនដំណឹងនេះ។ អ្នកប្រហែលជាត្រូវការបញ្ជាក់សមត្ថភាព ដល់កិច្ចការផ្ទៃក្នុងដូចជា ឆ្លើយតបនឹងសុំឱ្យអ្នកកាមរយ: Premera Blue Cross ។ ប្រសិនបើមានការសង្ស័យអ្វីមួយ អ្នកអាចទាក់ទងមន្ត្រីសេវាអតិថិជនរបស់យើង ឬប្រាកដថា យើងនឹងជួយអ្នកដោយមិនអស់លុយឡើយ។ សូមទូរស័ព្ទ 800-722-1471 (TTY: 800-842-5357)។

ਪੰਜਾਬੀ (Punjabi):

ਇਸ ਨੋਟਿਸ ਵਿਚ ਖਾਸ ਜਾਣਕਾਰੀ ਹੈ. ਇਸ ਨੋਟਿਸ ਵਿਚ Premera Blue Cross ਵਲੋਂ ਤੁਹਾਡੀ ਕਵਰੇਜ ਅਤੇ ਅਰਜੀ ਬਾਰੇ ਮਹੱਤਵਪੂਰਨ ਜਾਣਕਾਰੀ ਹੋ ਸਕਦੀ ਹੈ . ਇਸ ਨੋਟਿਸ ਨਵ ਖਾਸ ਤਾਰੀਖਾਂ ਹੋ ਸਕਦੀਆਂ ਹਨ. ਜੇਕਰ ਤੁਸੀਂ ਜਸਰਤ ਕਵਰੇਜ ਰਿੱਖਣੀ ਹੋਵੇ ਜਾਂ ਓਸ ਦੀ ਲਾਗਤ ਜਵਿੱਚ ਮਦਦ ਦੇ ਇਕੱਠ ਹੋ ਤਾਂ ਤੁਹਾਨੂੰ ਅੰਤਮ ਤਾਰੀਖ ਤੋਂ ਪਹਿਲਾਂ ਢੁੱਝ ਖਾਸ ਕਰਮ ਚੁੱਕਣ ਦੀ ਲੋੜ ਹੋ ਸਕਦੀ ਹੈ ,ਤੁਹਾਨੂੰ ਮੁਫਤ ਵਿੱਚ ਤੋਂ ਅਪਣੀ ਭਾਸ਼ਾ ਵਿੱਚ ਜਾਣਕਾਰੀ ਅਤੇ ਮਦਦ ਪ੍ਰਾਪਤ ਕਰਨ ਦਾ ਅਧਿਕਾਰ ਹੈ ,ਕਾਲ 800-722-1471 (TTY: 800-842-5357).

فارسی (Farsi):

این اعلامیه حاوی اطلاعات مهم میباشد. این اعلامیه ممکن است حاوی اطلاعات مهم درباره فرم تقاضا و یا پوشش بیمه ای شما از طریق Premera Blue Cross باشد. به تاریخ های مهم در این اعلامیه توجه نمایید. شما ممکن است برای حفظ پوشش بیمه تان یا کمک در پرداخت هزینه های درمانی تان، به تاریخ های مشخصی برای انجام کارهای خاصی احتیاج داشته باشید. شما حق این را دارید که این اطلاعات و کمک را به زبان خود به طور رایگان دریافت نمایید. برای کسب اطلاعات با شماره 800-722-1471 (کلیران TTY تماس باشماره 800-842-5357) تماس برقرار نمایید.

Polskie (Polish):

To ogłoszenie może zawierać ważne informacje. To ogłoszenie może zawierać ważne informacje odnośnie Państwa wniosku lub zakresu świadczeń poprzez Premera Blue Cross. Prosimy zwrócić uwagę na kluczowe daty, które mogą być zawarte w tym ogłoszeniu aby nie przekroczyć terminów w przypadku utrzymania polisy ubezpieczeniowej lub pomocy związanej z kosztami. Macie Państwo prawo do bezpłatnej informacji we własnym języku. Zadzwońcie pod 800-722-1471 (TTY: 800-842-5357).

Português (Portuguese):

Este aviso contém informações importantes. Este aviso poderá conter informações importantes a respeito de sua aplicação ou cobertura por meio do Premera Blue Cross. Poderão existir datas importantes neste aviso. Talvez seja necessário que você tome providências dentro de determinados prazos para manter sua cobertura de saúde ou ajuda de custos. Você tem o direito de obter esta informação e ajuda em seu idioma e sem custos. Ligue para 800-722-1471 (TTY: 800-842-5357).

Română (Romanian):

Prezenta notificare conține informații importante. Această notificare poate conține informații importante privind cererea sau acoperirea asigurării dumneavoastră de sănătate prin Premera Blue Cross. Pot exista date cheie în această notificare. Este posibil să fie nevoie să acționați până la anumite termene limită pentru a vă menține acoperirea asigurării de sănătate sau asistența provizorie la costuri. Aveți dreptul de a obține gratuit aceste informații și ajutor în limba dumneavoastră. Sunați la 800-722-1471 (TTY: 800-842-5357).

Русский (Russian):

Настоящее уведомление содержит важную информацию. Это уведомление может содержать важную информацию о вашем заявлении или страховом покрытии через Premera Blue Cross. В настоящем уведомлении могут быть указаны ключевые даты. Вам, возможно, потребуется принять меры к определенным предельным срокам для сохранения страхового покрытия или помощи с расходами. Вы имеете право на бесплатное получение этой информации и помощь на вашем языке. Звоните по телефону 800-722-1471 (TTY: 800-842-5357).

Fa'asamoa (Samoan):

Atonu ua iai i lenei fa'asilasilaga ni fa'amatalaga e sili ona taua e tatau ona e malamalama i ai. O lenei fa'asilasilaga o se fesoasoani e fa'amatala atili i ai i le tulaga o le polokalame, Premera Blue Cross, ua e tau fia maua atu i ai. Fa'amolemole, ia e iloilo fa'alelei i aso fa'apitoa olo'o iai i lenei fa'asilasilaga taua. Masalo o le'a iai ni feau e tatau ona e faia ao le'i aulia le aso ua ta'ua i lenei fa'asilasilaga ina ia e iai pea ma maua fesoasoani mai ai i le polokalame a le Malo olo'o e iai i ai. Olo'o iai iate oe le aia tatau e maua atu i lenei fa'asilasilaga ma lenei fa'matalaga i legagana e te malamalama i ai aunoa ma se togiga tupe. Vili atu i le telefoni 800-722-1471 (TTY: 800-842-5357).

Español (Spanish):

Este Aviso contiene información importante. Es posible que este aviso contenga información importante acerca de su solicitud o cobertura a través de Premera Blue Cross. Es posible que haya fechas clave en este aviso. Es posible que deba tomar alguna medida antes de determinadas fechas para mantener su cobertura médica o ayuda con los costos. Usted tiene derecho a recibir esta información y ayuda en su idioma sin costo alguno. Llame al 800-722-1471 (TTY: 800-842-5357).

Tagalog (Tagalog):

Ang Paunawa na ito ay naglalaman ng mahalagang impormasyon tungkol sa iyong aplikasyon o pagsakop sa pamamagitan ng Premera Blue Cross. Maaaring may mga mahalagang petsa dito sa paunawa. Maaring mangailangan ka na magsagawa ng hakbang sa ilang mga itinakdang panahon upang mapanatili ang iyong pagsakop sa kalusugan o tulong na walang gastos. May karapatan ka na makakuha ng ganiitong impormasyon at tulong sa iyong wika ng walang gastos. Tumawag sa 800-722-1471 (TTY: 800-842-5357).

ไทย (Thai):

ประกาศนี้มีข้อมูลสำคัญ ประกาศนี้อาจมีข้อมูลที่สำคัญเกี่ยวกับกาการสมัครหรือขอบเขตประกันสุขภาพของคุณผ่าน Premera Blue Cross และอาจมีกำหนดการในประกาศนี้ คุณอาจจะต้องดำเนินการภายในกำหนดระยะเวลาที่แน่นอนเพื่อจะรักษาการประกันสุขภาพของคุณหรือการช่วยเหลือที่มีค่าใช้จ่าย คุณมีสิทธิที่จะได้รับข้อมูลและความช่วยเหลือในภาษาของคุณโดยไม่มีค่าใช้จ่าย โทร 800-722-1471 (TTY: 800-842-5357)

Український (Ukrainian):

Це повідомлення містить важливу інформацію. Це повідомлення може містити важливу інформацію про Ваше звернення щодо страховального покриття через Premera Blue Cross. Зверніть увагу на ключові дати, які можуть бути вказані у цьому повідомленні. Існує імовірність того, що Вам треба буде здійснити певні кроки у конкретні кінцеві строки для того, щоб зберегти Ваше медичне страхування або отримати фінансову допомогу. У Вас є право на отримання цієї інформації та допомоги безкоштовно на Вашій рідній мові. Дзвоніть за номером телефону 800-722-1471 (TTY: 800-842-5357).

Tiếng Việt (Vietnamese):

Thông báo này cung cấp thông tin quan trọng. Thông báo này có thông tin quan trọng về đơn xin tham gia hoặc hợp đồng bảo hiểm của quý vị qua chương trình Premera Blue Cross. Xin xem ngày quan trọng trong thông báo này. Quý vị có thể phải thực hiện theo thông báo đúng trong thời hạn để duy trì bảo hiểm sức khỏe hoặc được trợ giúp thêm về chi phí. Quý vị có quyền được biết thông tin này và được trợ giúp bằng ngôn ngữ của mình miễn phí. Xin gọi số 800-722-1471 (TTY: 800-842-5357).