

5.70.58

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Subsection:	Analgesics and Anesthetics	Original Policy Date:	December 18, 2016
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Last Review Date: March 12, 2021

Butalbital Analgesics

Description

Allzital (butalbital-acetaminophen), butalbital-aspirin-caffeine, butalbital-aspirin-caffeine-codeine, butalbital-acetaminophen, butalbital-acetaminophen-caffeine, butalbital-acetaminophen-caffeine-codeine, Vanatol LQ (butalbital-acetaminophen-caffeine liquid oral solution)

Background

Butalbital containing products are non-opioid analgesics that contain a combination of different drug products indicated for the relief of the symptom complex of tension (or muscle contraction) headache pain. Butalbital is a short to intermediate-acting barbiturate that works in concert with acetaminophen, an antipyretic non-salicylate agent, aspirin, a pain-relieving NSAID, and caffeine, a stimulant that works in the CNS, to decrease pain via a mechanism that isn't well understood. Butalbital is a habit-forming drug that potentiates the effects of other commonly abuse drugs or substances like alcohol. Caffeine might help increase vasodilation and smooth muscle relaxation, while butalbital is thought to help balance the CNS stimulation caused by caffeine and produces depressant effects (1).

Regulatory Status

FDA approved indication: Butalbital containing products are used in the relief of the symptom complex of tension or muscle contraction headaches (2-8).

Frequent use of acute medications is generally thought to cause medication-overuse headache. To decrease the risk of medication-overuse headache ("rebound headache" or "drug-induced headache") many experts limit acute therapy to two headache days per week on a regular basis. Based on concerns of overuse, medication-overuse headache, and withdrawal, the use of

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butalbital-containing analgesics should be limited and carefully monitored. The Allzital limit is set to the maximum of 12 doses per day for acute treatment of 8 headaches per month as this product contains less butalbital than other products. The quantity limit for all other butalbital products is set to the maximum of 6 doses per day for acute treatment of 8 headaches per month (7).

Acetaminophen has been associated with cases of acute liver failure, at times resulting in liver transplant. Most of the cases of liver injury are associated with the use of acetaminophen at doses that exceed 4000 milligrams per day, and often involve more than one acetaminophen-containing product. Instruct patients to look for acetaminophen or APAP on package labels and not to use more than one product that contains acetaminophen. (5-6).

The FDA has determined that a REMS is necessary for all opioid analgesics intended for outpatient use to ensure that the benefits of these drugs continue to outweigh the risks. The Opioid Analgesic REMS is a strategy to reduce the risk of abuse, misuse, addiction, overdose, and deaths due to prescription opioid analgesics (9).

Safety and effectiveness of Allzital, butalbital-acetaminophen, butalbital-acetaminophen-caffeine, and Vanatol LQ in patients below the age of 12 have not been established. Safety and effectiveness of butalbital-acetaminophen-caffeine-codeine, butalbital-aspirin-caffeine, and butalbital-aspirin-caffeine-codeine in patients below the age of 18 have not been established (2-8).

Related policies

Amerge, Axert, Frova, Maxalt, Migraine CGRP Antagonists IV, Migraine CGRP Antagonists SC, Migraine CGRP Antagonists Oral, Migraine Powders, Migranal Nasal Spray, Relpax, Reyvow, Sumatriptan, Sumatriptan Injection, Zomig

Policy

This policy statement applies to clinical review performed for pre-service (Prior Approval, Precertification, Advanced Benefit Determination, etc.) and/or post-service claims.

Butalbital containing analgesics may be considered **medically necessary** in a patient suffering from tension or muscle contraction headaches if the conditions indicated below are met.

Butalbital containing analgesics may be considered **investigational use** for all other indications.

Prior-Approval Requirements

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Diagnosis

Patient must have the following:

1. Tension or muscle contraction headaches

Allzital, butalbital-acetaminophen and butalbital-acetaminophen-caffeine

AND ALL of the following:

- a. 12 years of age or older
- b. **NO** previous or current liver function concerns or cirrhosis
- c. Prescriber agrees to counsel regarding concurrent use of a product(s) containing acetaminophen

Vanatol LQ

AND ALL of the following:

- a. 12 years of age or older
- b. Inadequate response to generic butalbital-containing products
- c. **NO** previous or current liver function concerns or cirrhosis
- d. Prescriber agrees to counsel regarding concurrent use of a product(s) containing acetaminophen

Butalbital-acetaminophen-caffeine-codeine, butalbital-aspirin-caffeine, and butalbital-aspirin-caffeine-codeine

AND ALL of the following:

- a. 18 years of age or older
- b. **NO** previous or current liver function concerns or cirrhosis
- c. If acetaminophen containing product – prescriber agrees to counsel regarding concurrent use of a product(s) containing acetaminophen
- d. If codeine containing product – **NO** dual therapy with an opioid addiction therapy or methadone
- e. If codeine containing product – prescriber agrees to participate in the Opioid Analgesic REMS program and to monitor for abuse, misuse, addiction, and overdose and discontinue if necessary (<http://www.er-la-opioidrems.com/lwgUI/rems/home.action>)
- f. If codeine containing product – **NO** cumulative morphine milligram equivalent (MME) over 300 MME

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Prior – Approval *Renewal* Requirements

Same as above

Policy Guidelines

Pre - PA Allowance

Quantity

AGE 12 YEARS OF AGE AND OLDER	
Allzital (butalbital 25 mg / APAP 325 mg)	144 tabs per 90 days OR
Bupap (butalbital 50mg / APAP 300mg)	
Tencon (butalbital 50mg / APAP 325mg)	
Fioricet (butalbital 50mg / APAP 300mg / caffeine 40mg)	
Esgic (butalbital 50mg / APAP 325mg / caffeine 40mg)	
Esgic Plus (butalbital 50mg / APAP 500mg / caffeine 40mg)	
Vanatol LQ (butalbital / APAP / caffeine solution)	2,160 mL per 90 days

OR

AGE 18 YEARS OF AGE AND OLDER	
Allzital (butalbital 25 mg / APAP 325 mg)	144 tabs per 90 days OR
Bupap (butalbital 50mg / APAP 300mg)	
Tencon (butalbital 50mg / APAP 325mg)	
Fioricet (butalbital 50mg / APAP 300mg / caffeine 40mg)	
Esgic (butalbital 50mg / APAP 325mg / caffeine 40mg)	
Esgic Plus (butalbital 50mg / APAP 500mg / caffeine 40mg)	
Fiorinal (butalbital 50mg / aspirin 325mg / caffeine 40mg)	
Fiorinal with Codeine* (butalbital / aspirin / caffeine / codeine)	
Fioricet with Codeine* (butalbital / APAP/ caffeine / codeine)	2,160 mL per 90 days
Vanatol LQ (butalbital / APAP/ caffeine solution)	

*Patients are only eligible for a combination of opioids not exceeding 300 MME

Prior - Approval Limits

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Quantity

Allzital (butalbital 25 mg / APAP 325 mg)	288 tabs per 90 days OR
Bupap (butalbital 50mg / APAP 300mg)	216 tabs per 90 days OR
Tencon (butalbital 50mg / APAP 325mg)	216 tabs per 90 days OR
Fioricet (butalbital 50mg / APAP 300mg / caffeine 40mg)	216 tabs per 90 days OR
Esgic (butalbital 50mg / APAP 325mg / caffeine 40mg)	216 tabs per 90 days OR
Esgic Plus (butalbital 50mg / APAP 500mg / caffeine 40mg)	216 tabs per 90 days OR
Fiorinal (butalbital 50mg / aspirin 325mg / caffeine 40mg)	216 tabs per 90 days OR
Fiorinal with Codeine (butalbital / aspirin / caffeine / codeine)	216 tabs per 90 days OR
Fioricet with Codeine (butalbital / APAP / caffeine / codeine)	216 tabs per 90 days OR
Vanatol LQ (butalbital / APAP / caffeine solution)	3,240 mL per 90 days

Duration 6 months

Prior – Approval *Renewal* Limits

Same as above

Rationale**Summary**

Butalbital is a short to intermediate-acting barbiturate that causes CNS depression. Caffeine is a CNS stimulant that is thought to help increase vasodilation (smooth muscle relaxation). Acetaminophen might help decrease pain sensation in the peripheral nervous system by blocking those signals. Aspirin is an NSAID that decreases pain and swelling by blocking prostaglandins. Butalbital-containing analgesics are FDA approved for the treatment of the symptom complex of tension or muscle contraction headache. These products can have pronounced sedative effects and butalbital is habit-forming (1-8).

Prior authorization is required to ensure the safe, clinically appropriate and cost-effective use of butalbital-containing analgesics, while maintaining optimal therapeutic outcomes.

References

1. Bryczkowski, C., Geib, A.J. Combined Butalbital/Acetaminophen/Caffeine Overdose: Case Files of the Robert Wood Johnson Medical School Toxicology Service. Journal of Medical Toxicology, Sept. 26, 2012.
2. Vanatol LQ [package insert]. Arlington, TX GM Pharmaceuticals, Inc.; November 2019.

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3. Fiorinal [package insert]. Parsippany, NJ: Actavis Pharma, Inc.; November 2018.
4. Fiorinal with Codeine [package insert]. Madison, NJ: Allergan USA, Inc.; October 2019.
5. Fioricet [package insert]. Parsippany, NJ: Actavis Pharma, Inc.; June 2020.
6. Fioricet with Codeine [package insert]. Parsippany, NJ: Actavis Pharma, Inc.; February 2020.
7. Allzital [package insert]. Canton, MS: Skylar Laboratories, LLC.; March 2019.
8. Beithon J, Gallenberg M, Johnson K, et al. Institute for Clinical Systems Improvement. Diagnosis and Treatment of Headache. https://www.icsi.org/guidelines__more/catalog_guidelines_and_more/catalog_guidelines/catalog_neurological_guidelines/headache. Updated January 2013.
9. Opioid Analgesic Risk Evaluation and Mitigation Strategy (REMS). Drug Safety and Availability: U.S. Food & Drug Administration. September 19, 2018.

Policy History

Date	Action
December 2016	New addition to PA
March 2017	Addition of age requirements to the Pre – PA Allowance
June 2017	Annual review
June 2017	Addition of no dual therapy with methadone
September 2017	Annual review
March 2018	Annual editorial review and reference update
October 2018	Addition of Opioid Analgesic REMS requirement to codeine-containing products
November 2018	Annual review and reference update. Addition of Opioid Analgesic REMS link per SME
March 2019	Annual review
December 2019	Annual review. Addition of requirement of no cumulative MME over 300. Revised requirement for prescriber to agree to counsel regarding using concurrent APAP medications
March 2020	Annual review and reference update
June 2020	Annual review
March 2021	Annual review and reference update

Keywords

This policy was approved by the FEP® Pharmacy and Medical Policy Committee on March 12, 2021 and is effective on April 1, 2021.