

Updates to the Alberta Drug Benefit List

Effective December 1, 2014



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Administered by Alberta Blue Cross
on behalf of Alberta Health.

The Drug Benefit List (DBL) is a list of drugs for which coverage may be provided to program participants. The DBL is not intended to be, and must not be used as a diagnostic or prescribing tool. Inclusion of a drug on the DBL does not mean or imply that the drug is fit or effective for any specific purpose. Prescribing professionals must always use their professional judgment and should refer to product monographs and any applicable practice guidelines when prescribing drugs. The product monograph contains information that may be required for the safe and effective use of the product.

Copies of the *Alberta Drug Benefit List* Publication are available from Pharmacy Services, Alberta Blue Cross at the address shown above.

Binder and contents: **\$42.00** (\$40.00 + \$2.00 G.S.T.)
Contents only: **\$36.75** (\$35.00 + \$1.75 G.S.T.)

A cheque or money order must accompany the request for copies.

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Special Authorization

The following drug product(s) will be considered for coverage by special authorization for patients covered under Alberta government-sponsored drug programs. Criteria for coverage of Alberta Human Services can be found in the December 1, 2014 Updates To the Alberta Human Services Drug Benefit Supplement.

New Drug Product(s) Available by Special Authorization

Trade Name / Strength / Form	Generic Description	DIN	MFR
AUBAGIO 14 MG TABLET	TERIFLUNOMIDE	00002416328	GZM

Additional Brand(s) and/or Strength(s) of Drug Products Available by Special Authorization

Trade Name / Strength / Form	Generic Description	DIN	MFR
APO-EZETIMIBE 10 MG TABLET	EZETIMIBE	00002427826	APX
EZETIMIBE 10 MG TABLET	EZETIMIBE	00002429659	SIV
EZETIMIBE 10 MG TABLET	EZETIMIBE	00002431300	SNS
JAMP-DONEPEZIL 5 MG TABLET	DONEPEZIL HCL	00002416948	JPC
JAMP-DONEPEZIL 10 MG TABLET	DONEPEZIL HCL	00002416956	JPC
MED-DUTASTERIDE 0.5 MG CAPSULE	DUTASTERIDE	00002416298	GMP

New Drug Product(s) Available by Step Therapy / Special Authorization

Trade Name / Strength / Form	Generic Description	DIN	MFR
JENTADUETO 2.5 MG / 500 MG TABLET	LINAGLIPTIN/ METFORMIN HCL	00002403250	BOE
JENTADUETO 2.5 MG / 850 MG TABLET	LINAGLIPTIN/ METFORMIN HCL	00002403269	BOE
JENTADUETO 2.5 MG / 1,000 MG TABLET	LINAGLIPTIN/ METFORMIN HCL	00002403277	BOE

Additional Brand(s) and/or Strength(s) of Drug Products Available by Restricted Benefit / Special Authorization

Trade Name / Strength / Form	Generic Description	DIN	MFR
JAMP-RIZATRIPTAN IR 5 MG TABLET	RIZATRIPTAN BENZOATE	00002429233	JPC
JAMP-RIZATRIPTAN IR 10 MG TABLET	RIZATRIPTAN BENZOATE	00002429241	JPC

Added Product(s)

Trade Name / Strength / Form	Generic Description	DIN	MFR
ABBOTT-RABEPRAZOLE 10 MG ENTERIC-COATED TABLET	RABEPRAZOLE SODIUM	00002422638	ABB
AMLODIPINE 5 MG TABLET	AMLODIPINE BESYLATE	00002429217	JPC
AMLODIPINE 10 MG TABLET	AMLODIPINE BESYLATE	00002429225	JPC
CALCITRIOL-ODAN 0.25 MCG CAPSULE	CALCITRIOL	00002431637	ODN
CALCITRIOL-ODAN 0.5 MCG CAPSULE	CALCITRIOL	00002431645	ODN
CITALOPRAM 20 MG TABLET	CITALOPRAM HYDROBROMIDE	00002430541	JPC
CITALOPRAM 40 MG TABLET	CITALOPRAM HYDROBROMIDE	00002430568	JPC
PMS-LATANOPROST 0.005 % OPHTHALMIC SOLUTION	LATANOPROST	00002317125	PMS

New Established Interchangeable (IC) Grouping(s)

The following IC Grouping(s) have been established and LCA pricing will be applied effective January 1, 2015.

Generic Description	Strength / Form	New LCA Price
CALCITRIOL	0.25 MCG CAPSULE	0.6960
CALCITRIOL	0.5 MCG CAPSULE	1.1069

Product(s) With A Price Change

The following product(s) had a Price Decrease. The previous higher price will be recognized until December 31, 2014.

Trade Name / Strength / Form	Generic Description	DIN	MFR
JAMP-RIZATRIPTAN 5 MG TABLET	RIZATRIPTAN BENZOATE	00002380455	JPC

Discontinued Listing(s)

Notification of discontinuation has been received from the manufacturer(s). The Alberta government-sponsored drug programs previously covered the following drug product(s). Effective December 1, 2014, the listed product(s) will no longer be a benefit and will not be considered for coverage by special authorization. A transition period will be applied and, as of December 31, 2014 claims will no longer pay for these products. Please note, for products that were covered by Special Authorization, no transition period will be applied, and as of November 30, 2014, claims will no longer pay for these products.

Trade Name / Strength / Form	Generic Description	DIN	MFR
NERISONE 0.1 % TOPICAL CREAM	DIFLUCORTOLONE VALERATE	00000587826	GSK
SARNA HC 1 % TOPICAL LOTION	HYDROCORTISONE	00000578541	GSK

Discontinued Listing(s), continued

Trade Name / Strength / Form	Generic Description	DIN	MFR
SARNA HC 2.5 % TOPICAL LOTION	HYDROCORTISONE	00000856711	GSK
TEVA-ATENOLOL 50 MG TABLET	ATENOLOL	00001912062	TEV
TEVA-ATENOLOL 100 MG TABLET	ATENOLOL	00001912054	TEV
TEVA-FLUTAMIDE 250 MG TABLET	FLUTAMIDE	00002230089	TEV
ZOVIRAX 200 MG TABLET	ACYCLOVIR	00000634506	GSK

PART 2

Drug Additions

ALBERTA DRUG BENEFIT LIST UPDATE

AMLODIPINE BESYLATE

5 MG (BASE) ORAL TABLET				
00002331284	AMLODIPINE	SNS	\$	0.2417
00002331934	AMLODIPINE	RAN	\$	0.2417
00002385791	AMLODIPINE	SIV	\$	0.2417
00002429217	AMLODIPINE	JPC	\$	0.2417
00002378760	AMLODIPINE-ODAN	ODN	\$	0.2417
00002273373	APO-AMLODIPINE	APX	\$	0.2417
00002397072	AURO-AMLODIPINE	AUR	\$	0.2417
00002297485	CO AMLODIPINE	APH	\$	0.2417
00002280132	GD-AMLODIPINE	GMD	\$	0.2417
00002331071	JAMP-AMLODIPINE	JPC	\$	0.2417
00002357194	JAMP-AMLODIPINE	JPC	\$	0.2417
00002371715	MAR-AMLODIPINE	MAR	\$	0.2417
00002362651	MINT-AMLODIPINE	MPI	\$	0.2417
00002272113	MYLAN-AMLODIPINE	MYP	\$	0.2417
00002326779	PHL-AMLODIPINE	PHH	\$	0.2417
00002284065	PMS-AMLODIPINE	PMS	\$	0.2417
00002321858	RAN-AMLODIPINE	RAN	\$	0.2417
00002259605	RATIO-AMLODIPINE	RPH	\$	0.2417
00002284383	SANDOZ AMLODIPINE	SDZ	\$	0.2417
00002357712	SEPTA-AMLODIPINE	SEP	\$	0.2417
00002250497	TEVA-AMLODIPINE	TEV	\$	0.2417
00000878928	NORVASC	PFI	\$	1.3426
10 MG (BASE) ORAL TABLET				
00002331292	AMLODIPINE	SNS	\$	0.3587
00002331942	AMLODIPINE	RAN	\$	0.3587
00002385805	AMLODIPINE	SIV	\$	0.3587
00002429225	AMLODIPINE	JPC	\$	0.3587
00002378779	AMLODIPINE-ODAN	ODN	\$	0.3587
00002273381	APO-AMLODIPINE	APX	\$	0.3587
00002397080	AURO-AMLODIPINE	AUR	\$	0.3587
00002297493	CO AMLODIPINE	APH	\$	0.3587
00002280140	GD-AMLODIPINE	GMD	\$	0.3587
00002331098	JAMP-AMLODIPINE	JPC	\$	0.3587
00002357208	JAMP-AMLODIPINE	JPC	\$	0.3587
00002371723	MAR-AMLODIPINE	MAR	\$	0.3587
00002362678	MINT-AMLODIPINE	MPI	\$	0.3587
00002272121	MYLAN-AMLODIPINE	MYP	\$	0.3587
00002326787	PHL-AMLODIPINE	PHH	\$	0.3587
00002284073	PMS-AMLODIPINE	PMS	\$	0.3587
00002321866	RAN-AMLODIPINE	RAN	\$	0.3587
00002259613	RATIO-AMLODIPINE	RPH	\$	0.3587
00002284391	SANDOZ AMLODIPINE	SDZ	\$	0.3587
00002357720	SEPTA-AMLODIPINE	SEP	\$	0.3587
00002250500	TEVA-AMLODIPINE	TEV	\$	0.3587
00000878936	NORVASC	PFI	\$	1.9930

CALCITRIOL

0.25 MCG ORAL CAPSULE				
00002431637	CALCITRIOL-ODAN	ODN	\$	0.6960
00000481823	ROCALTROL	HLR	\$	0.9280
0.5 MCG ORAL CAPSULE				
00002431645	CALCITRIOL-ODAN	ODN	\$	1.1069
00000481815	ROCALTROL	HLR	\$	1.4758

The DBL is not a prescribing or a diagnostic tool. Prescribers should refer to drug monographs and utilize professional judgment.

ALBERTA DRUG BENEFIT LIST UPDATE

CITALOPRAM HYDROBROMIDE

20 MG (BASE) ORAL TABLET

00002414589	ABBOTT-CITALOPRAM	ABB	\$	0.2397
00002355256	ACCEL-CITALOPRAM	ACP	\$	0.2397
00002246056	APO-CITALOPRAM	APX	\$	0.2397
00002275562	AURO-CITALOPRAM	AUR	\$	0.2397
00002331950	CITALOPRAM	RAN	\$	0.2397
00002353660	CITALOPRAM	SNS	\$	0.2397
00002387956	CITALOPRAM	SIV	\$	0.2397
00002430541	CITALOPRAM	JPC	\$	0.2397
00002306239	CITALOPRAM-ODAN	ODN	\$	0.2397
00002248050	CO CITALOPRAM	APH	\$	0.2397
00002313405	JAMP-CITALOPRAM	JPC	\$	0.2397
00002371898	MAR-CITALOPRAM	MAR	\$	0.2397
00002304686	MINT-CITALOPRAM	MPI	\$	0.2397
00002246594	MYLAN-CITALOPRAM	MYP	\$	0.2397
00002409011	NAT-CITALOPRAM	NTP	\$	0.2397
00002248944	PHL-CITALOPRAM	PHH	\$	0.2397
00002248010	PMS-CITALOPRAM	PMS	\$	0.2397
00002285622	RAN-CITALO	RAN	\$	0.2397
00002252112	RATIO-CITALOPRAM	RPH	\$	0.2397
00002248170	SANDOZ CITALOPRAM	SDZ	\$	0.2397
00002355272	SEPTA-CITALOPRAM	SEP	\$	0.2397
00002293218	TEVA-CITALOPRAM	TEV	\$	0.2397
00002239607	CELEXA	LBC	\$	1.3437

40 MG (BASE) ORAL TABLET

00002414597	ABBOTT-CITALOPRAM	ABB	\$	0.2397
00002355264	ACCEL-CITALOPRAM	ACP	\$	0.2397
00002246057	APO-CITALOPRAM	APX	\$	0.2397
00002275570	AURO-CITALOPRAM	AUR	\$	0.2397
00002331977	CITALOPRAM	RAN	\$	0.2397
00002353679	CITALOPRAM	SNS	\$	0.2397
00002387964	CITALOPRAM	SIV	\$	0.2397
00002430568	CITALOPRAM	JPC	\$	0.2397
00002306247	CITALOPRAM-ODAN	ODN	\$	0.2397
00002248051	CO CITALOPRAM	APH	\$	0.2397
00002313413	JAMP-CITALOPRAM	JPC	\$	0.2397
00002371901	MAR-CITALOPRAM	MAR	\$	0.2397
00002304694	MINT-CITALOPRAM	MPI	\$	0.2397
00002246595	MYLAN-CITALOPRAM	MYP	\$	0.2397
00002409038	NAT-CITALOPRAM	NTP	\$	0.2397
00002248945	PHL-CITALOPRAM	PHH	\$	0.2397
00002248011	PMS-CITALOPRAM	PMS	\$	0.2397
00002285630	RAN-CITALO	RAN	\$	0.2397
00002252120	RATIO-CITALOPRAM	RPH	\$	0.2397
00002248171	SANDOZ CITALOPRAM	SDZ	\$	0.2397
00002355280	SEPTA-CITALOPRAM	SEP	\$	0.2397
00002293226	TEVA-CITALOPRAM	TEV	\$	0.2397
00002239608	CELEXA	LBC	\$	1.3437

LATANOPROST

0.005 % OPHTHALMIC SOLUTION

00002296527	APO-LATANOPROST	APX	\$	3.6320
00002254786	CO LATANOPROST	APH	\$	3.6320
00002373041	GD-LATANOPROST	GMD	\$	3.6320
00002317125	PMS-LATANOPROST	PMS	\$	3.6320
00002367335	SANDOZ LATANOPROST	SDZ	\$	3.6320
00002231493	XALATAN	PFI	\$	11.1758

The DBL is not a prescribing or a diagnostic tool. Prescribers should refer to drug monographs and utilize professional judgment.

RABEPRAZOLE SODIUM

10 MG ORAL ENTERIC-COATED TABLET

00002422638	ABBOTT-RABEPRAZOLE	ABB	\$	0.1204
00002345579	APO-RABEPRAZOLE	APX	\$	0.1204
00002408392	MYLAN-RABEPRAZOLE	MYP	\$	0.1204
00002310805	PMS-RABEPRAZOLE EC	PMS	\$	0.1204
00002385449	RABEPRAZOLE	SIV	\$	0.1204
00002356511	RABEPRAZOLE EC	SNS	\$	0.1204
00002298074	RAN-RABEPRAZOLE	RAN	\$	0.1204
00002314177	SANDOZ RABEPRAZOLE	SDZ	\$	0.1204
00002296632	TEVA-RABEPRAZOLE	TEV	\$	0.1204
00002243796	PARIET	JAI	\$	0.6688

RIZATRIPTAN BENZOATE

RESTRICTED BENEFIT - This product is a benefit for patients 18 to 64 years of age inclusive for the treatment of acute migraine attacks in patients where standard therapy has failed. (Refer to Criteria for Special Authorization of Select Drug Products of the List for eligibility in patients 65 years of age and older; and Criteria for Special Authorization of Select Drug Products of the Alberta Human Services Drug Benefit Supplement for eligibility in Alberta Human Services clients.)

5 MG (BASE) ORAL TABLET

00002393468	APO-RIZATRIPTAN	APX	\$	3.7050
00002380455	JAMP-RIZATRIPTAN	JPC	\$	3.7050
00002429233	JAMP-RIZATRIPTAN IR	JPC	\$	3.7050

10 MG (BASE) ORAL TABLET

00002393476	APO-RIZATRIPTAN	APX	\$	3.7050
00002381702	CO RIZATRIPTAN	APH	\$	3.7050
00002380463	JAMP-RIZATRIPTAN	JPC	\$	3.7050
00002429241	JAMP-RIZATRIPTAN IR	JPC	\$	3.7050
00002379678	MAR-RIZATRIPTAN	MAR	\$	3.7050
00002240521	MAXALT	MFC	\$	15.5874

PART 3

Special Authorization

**ALBERTA DRUG BENEFIT LIST UPDATE
CRITERIA FOR SPECIAL AUTHORIZATION OF SELECT DRUG PRODUCTS**

DONEPEZIL HCL

"For the treatment of Alzheimer's disease in patients with an MMSE (Mini Mental State Exam) score between 10-26 and/or an InterRAI-Cognitive Performance Scale score between 1-4.

Coverage cannot be provided for two or more medications used in the treatment of Alzheimer's disease (donepezil, galantamine, rivastigmine) when these medications are intended for use in combination.

Special authorization coverage may be granted for a maximum of 24 months per request.

For each request, an updated MMSE score or InterRAI-Cognitive Performance Scale score and the date on which the exam was administered must be provided.

Renewal requests may be considered for patients where the updated MMSE score is 10 or higher or the InterRAI-Cognitive Performance Scale is 4 or lower while on this drug."

All requests (including renewal requests) for donepezil HCl must be completed using the Donepezil/Galantamine/Rivastigmine Special Authorization Request Form (ABC 30776).

5 MG ORAL TABLET

00002362260	APO-DONEPEZIL	APX	\$	1.1806
00002400561	AURO-DONEPEZIL	AUR	\$	1.1806
00002397595	CO DONEPEZIL	APH	\$	1.1806
00002420597	DONEPEZIL	SIV	\$	1.1806
00002402645	DONEPEZIL HYDROCHLORIDE	AHI	\$	1.1806
00002404419	JAMP-DONEPEZIL	JPC	\$	1.1806
00002416948	JAMP-DONEPEZIL	JPC	\$	1.1806
00002402092	MAR-DONEPEZIL	MAR	\$	1.1806
00002359472	MYLAN-DONEPEZIL	MYP	\$	1.1806
00002322331	PMS-DONEPEZIL	PMS	\$	1.1806
00002381508	RAN-DONEPEZIL	RAN	\$	1.1806
00002328666	SANDOZ DONEPEZIL	SDZ	\$	1.1806
00002428482	SEPTA DONEPEZIL	SEP	\$	1.1806
00002340607	TEVA-DONEPEZIL	TEV	\$	1.1806
00002232043	ARICEPT	PFI	\$	4.8620

10 MG ORAL TABLET

00002362279	APO-DONEPEZIL	APX	\$	1.1806
00002400588	AURO-DONEPEZIL	AUR	\$	1.1806
00002397609	CO DONEPEZIL	APH	\$	1.1806
00002420600	DONEPEZIL	SIV	\$	1.1806
00002402653	DONEPEZIL HYDROCHLORIDE	AHI	\$	1.1806
00002404427	JAMP-DONEPEZIL	JPC	\$	1.1806
00002416956	JAMP-DONEPEZIL	JPC	\$	1.1806
00002402106	MAR-DONEPEZIL	MAR	\$	1.1806
00002359480	MYLAN-DONEPEZIL	MYP	\$	1.1806
00002322358	PMS-DONEPEZIL	PMS	\$	1.1806
00002381516	RAN-DONEPEZIL	RAN	\$	1.1806
00002328682	SANDOZ DONEPEZIL	SDZ	\$	1.1806
00002428490	SEPTA DONEPEZIL	SEP	\$	1.1806
00002340615	TEVA-DONEPEZIL	TEV	\$	1.1806
00002232044	ARICEPT	PFI	\$	4.8620

**ALBERTA DRUG BENEFIT LIST UPDATE
CRITERIA FOR SPECIAL AUTHORIZATION OF SELECT DRUG PRODUCTS**

DUTASTERIDE

"For the treatment of benign prostatic hyperplasia in patients who are poor surgical risks or who have enlarged prostates and have moderate to severe symptoms suggestive of obstruction.

Special authorization may be granted for 6 months"

Information is required regarding the medical condition(s) or circumstances by which this patient would be deemed a poor surgical risk.

All requests (including renewal requests) for dutasteride must be completed using the Dutasteride/Finasteride Special Authorization Request Form (ABC 31257).

The following product(s) are eligible for auto-renewal.

0.5 MG ORAL CAPSULE

00002412691	ACT DUTASTERIDE	APH	\$	0.4205
00002404206	APO-DUTASTERIDE	APX	\$	0.4205
00002416298	MED-DUTASTERIDE	GMP	\$	0.4205
00002428873	MINT-DUTASTERIDE	MPI	\$	0.4205
00002393220	PMS-DUTASTERIDE	PMS	\$	0.4205
00002424444	SANDOZ DUTASTERIDE	SDZ	\$	0.4205
00002408287	TEVA-DUTASTERIDE	TEV	\$	0.4205
00002247813	AVODART	GSK	\$	1.6819

**ALBERTA DRUG BENEFIT LIST UPDATE
CRITERIA FOR SPECIAL AUTHORIZATION OF SELECT DRUG PRODUCTS**

EZETIMIBE

"For the treatment of hypercholesterolemia in patients who are intolerant to statins or in whom a statin is contraindicated and who are at high cardiovascular risk*; or

For the treatment of hypercholesterolemia when used in combination with a statin in patients failing to achieve target LDL with a statin at maximum tolerable dose or maximum recommended dose as per respective product monograph and who are at high cardiovascular risk*:

* High cardiovascular risk is defined as possessing one of the following:

- 1) Pre-existing cardiovascular disease and/or cerebrovascular disease, or
- 2) Diabetes, or
- 3) Familial hypercholesterolemia, or
- 4) Greater than or equal to 20% risk as defined by the Framingham Risk Assessment Tool, or
- 5) Three or more of the following risk factors:
 - Family history of premature cardiovascular disease
 - Smoking
 - Hypertension
 - Obesity
 - Glucose intolerance
 - Renal disease.

Special authorization for these criteria may be granted for 6 months."

All requests for ezetimibe must be completed using the Ezetimibe Special Authorization Request Form (ABC 30925).

The following product(s) are eligible for auto-renewal.

10 MG ORAL TABLET

00002414716	ACT EZETIMIBE	APH	\$	0.4549
00002427826	APO-EZETIMIBE	APX	\$	0.4549
00002429659	EZETIMIBE	SIV	\$	0.4549
00002431300	EZETIMIBE	SNS	\$	0.4549
00002378035	MYLAN-EZETIMIBE	MYP	\$	0.4549
00002419548	RAN-EZETIMIBE	RAN	\$	0.4549
00002416778	SANDOZ EZETIMIBE	SDZ	\$	0.4549
00002354101	TEVA-EZETIMIBE	TEV	\$	0.4549
00002247521	EZETROL	MFC	\$	1.8196

ALBERTA DRUG BENEFIT LIST UPDATE
CRITERIA FOR SPECIAL AUTHORIZATION OF SELECT DRUG PRODUCTS

LINAGLIPTIN/ METFORMIN HCL

SPECIAL AUTHORIZATION

The drug product(s) listed below are eligible for coverage via the step therapy/special authorization process.

FIRST-LINE DRUG PRODUCT(S): METFORMIN
SECOND-LINE DRUG PRODUCT(S): SULFONYLUREAS
AND WHERE INSULIN IS NOT AN OPTION

As add-on therapy for the treatment of Type 2 diabetes in patients with intolerance to and/or inadequate glycemic control on:

- a sufficient trial (i.e. a minimum of 6 months) of metformin, AND
- a sulfonylurea, AND
- for whom insulin is not an option.

Or, for whom these products are contraindicated.

Special authorization may be granted for 24 months.

Note: If a claim for the Step therapy drug product is rejected, pharmacists can use their professional judgment to determine the appropriateness of using the intervention code(s) noted below to re-submit a claim. The pharmacist is responsible to document on the patient's record the rationale for using the second-line therapy drug.

- UP - First-line therapy ineffective
- UQ - First-line therapy not tolerated
- CA - Prior adverse reaction
- CB - Previous treatment failure
- CJ - Product is not effective

All requests for linagliptin+metformin must be completed using the DPP-4 Inhibitors Special Authorization Request Form (ABC 60012).

2.5 MG * 500 MG ORAL TABLET			
00002403250 JENTADUETO	BOE	\$	1.3337
2.5 MG * 850 MG ORAL TABLET			
00002403269 JENTADUETO	BOE	\$	1.3337
2.5 MG * 1,000 MG ORAL TABLET			
00002403277 JENTADUETO	BOE	\$	1.3337

ALBERTA DRUG BENEFIT LIST UPDATE
CRITERIA FOR SPECIAL AUTHORIZATION OF SELECT DRUG PRODUCTS

RIZATRIPTAN BENZOATE

(Refer to 28:32.28 of the Alberta Drug Benefit List for coverage of patients 18 to 64 years of age inclusive.)

"For the treatment of acute migraine attacks in patients 65 years of age and older where other standard therapy has failed."

"For the treatment of acute migraine attacks in patients 65 years of age and older who have been using rizatriptan benzoate prior to turning 65."

"Special authorization for both criteria may be granted for 24 months."

In order to comply with the first criteria, information is required regarding previous medications utilized and the patient's response to therapy.

The following product(s) are eligible for auto-renewal.

5 MG (BASE) ORAL TABLET

00002393468	APO-RIZATRIPTAN	APX	\$	3.7050
00002380455	JAMP-RIZATRIPTAN	JPC	\$	3.7050
00002429233	JAMP-RIZATRIPTAN IR	JPC	\$	3.7050

10 MG (BASE) ORAL TABLET

00002393476	APO-RIZATRIPTAN	APX	\$	3.7050
00002381702	CO RIZATRIPTAN	APH	\$	3.7050
00002380463	JAMP-RIZATRIPTAN	JPC	\$	3.7050
00002429241	JAMP-RIZATRIPTAN IR	JPC	\$	3.7050
00002379678	MAR-RIZATRIPTAN	MAR	\$	3.7050
00002240521	MAXALT	MFC	\$	15.5874

**ALBERTA DRUG BENEFIT LIST UPDATE
CRITERIA FOR SPECIAL AUTHORIZATION OF SELECT DRUG PRODUCTS**

TERIFLUNOMIDE

SPECIAL AUTHORIZATION

Relapsing Remitting Multiple Sclerosis (RRMS):

Special authorization coverage may be provided for the reduction of the frequency and severity of clinical relapses and reduction of the number and volume of active brain lesions, identified on MRI scans, in ambulatory patients with relapsing remitting multiple sclerosis.

Coverage

For coverage, this drug must be prescribed by a registered MS Neurologist. A current assessment must be completed by a registered MS Neurologist at every request.

To register to become an MS Neurologist please complete the Registration for MS Neurologist Status Form (ABC 60002).

Initial Coverage

- 1) The registered MS Neurologist must confirm a diagnosis of RRMS;
- 2) The patient must have active disease which is defined as at least two relapses* of MS during the previous two years or in the two years prior to starting an MS disease modifying therapy (DMT).

*A relapse is defined as the appearance of new symptoms or worsening of old symptoms, lasting at least 48 hours in the absence of fever, not associated with withdrawal from steroids. Onset of clinical relapses must be separated by a period of at least one month. At least one definite gadolinium-enhancing T1 MRI lesion (not questionable faint enhancement) obtained at least 90 days after initiation of the DMT and at least 90 days before or after a relapse may substitute for one clinical relapse.

- 3) The patient must be ambulatory with or without aid (The registered MS Neurologist must provide a current updated Expanded Disability Status Scale (EDSS) score less than or equal to 6.5).

Coverage may be approved for up to 12 months. Patients will be limited to receiving a one-month supply of teriflunomide per prescription at their pharmacy for the first 12 months of coverage.

Continued Coverage

For continued coverage beyond the initial coverage period, the patient must meet the following criteria:

- 1) The patient must be assessed by a registered MS Neurologist;
- 2) The registered MS Neurologist must confirm a diagnosis of RRMS;
- 3) The registered MS Neurologist must provide a current updated EDSS score. The patient must not have an EDSS score of 7.0 or above sustained for one year or more.

Coverage of this drug may be considered in a patient with a sustained EDSS score of 7.0 or above in exceptional circumstances. For MS DMT coverage to be considered, details of the exceptional circumstance must be provided in a letter from the registered MS Neurologist and accompany the Special Authorization Request Form.

Continued coverage may be approved for up to 12 months. Patients may receive up to 100 days' supply of teriflunomide per prescription at their pharmacy.

ALBERTA DRUG BENEFIT LIST UPDATE
CRITERIA FOR SPECIAL AUTHORIZATION OF SELECT DRUG PRODUCTS

TERIFLUNOMIDE

Restarting After an Interruption in Therapy Greater Than 12 Months

In order to be eligible for coverage, after an interruption in therapy greater than 12 months, the patient must meet the following criteria:

- 1) At least one relapse* per 12 month period; or
- 2) At least two relapses* during the previous 24 month period.

All requests (including renewal requests) for teriflunomide must be completed using the Dimethyl Fumarate/Glatiramer Acetate/ Interferon Beta-1a/Interferon Beta-1b/Teriflunomide Special Authorization Request Form (ABC 60001).

14 MG ORAL TABLET

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