

SENATE HEALTH AND PUBLIC AFFAIRS COMMITTEE SUBSTITUTE FOR
SENATE BILL 135

56TH LEGISLATURE - STATE OF NEW MEXICO - SECOND SESSION, 2024

This document may incorporate amendments proposed by a committee, but not yet adopted, as well as amendments that have been adopted during the current legislative session. The document is a tool to show amendments in context and cannot be used for the purpose of adding amendments to legislation.

AN ACT

RELATING TO HEALTH COVERAGE; AMENDING THE HEALTH CARE PURCHASING ACT, THE PUBLIC ASSISTANCE ACT, THE NEW MEXICO INSURANCE CODE, THE HEALTH MAINTENANCE ORGANIZATION LAW AND THE NONPROFIT HEALTH CARE PLAN LAW TO MODIFY THE GUIDELINES RELATING TO STEP THERAPY FOR PRESCRIPTION DRUG COVERAGE AND ELIMINATE STEP THERAPY REQUIREMENTS FOR CERTAIN CONDITIONS.

BE IT ENACTED BY THE LEGISLATURE OF THE STATE OF NEW MEXICO:

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SECTION 1. Section 13-7-18 NMSA 1978 (being Laws 2018, Chapter 9, Section 1) is amended to read:

"13-7-18. PRESCRIPTION DRUG COVERAGE--STEP THERAPY PROTOCOLS--CLINICAL REVIEW CRITERIA--EXCEPTIONS.--

A. Group health coverage, including any form of self-insurance, offered, issued or renewed under the Health Care Purchasing Act that provides coverage for prescription drugs for which any step therapy protocols are required shall establish clinical review criteria for those step therapy protocols. The clinical review criteria shall be based on clinical practice guidelines that:

(1) recommend that the prescription drugs subject to step therapy protocols be taken in the specific sequence required by the step therapy protocol;

(2) are developed and endorsed by an interdisciplinary panel of experts that manages conflicts of interest among the members of the panel of experts by:

(a) requiring members to: 1) disclose any potential conflicts of interest with group health plan administrators, insurers, health maintenance organizations, health care plans, pharmaceutical manufacturers, pharmacy benefits managers and any other entities; and 2) recuse themselves if there is a conflict of interest; and

(b) using analytical and methodological experts to work to provide objectivity in data analysis and

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ranking of evidence through the preparation of evidence tables and facilitating consensus;

(3) are based on high-quality studies, research and medical practice;

(4) are created pursuant to an explicit and transparent process that:

(a) minimizes bias and conflicts of interest;

(b) explains the relationship between treatment options and outcomes;

(c) rates the quality of the evidence supporting recommendations; and

(d) considers relevant patient subgroups and preferences; and

(5) take into account the needs of atypical patient populations and diagnoses.

B. In the absence of clinical guidelines that meet the requirements of Subsection A of this section, peer-reviewed publications may be substituted.

C. When a group health plan restricts coverage of a prescription drug for the treatment of any medical condition through the use of a step therapy protocol, an enrollee and the practitioner prescribing the prescription drug shall have access to a clear, readily accessible and convenient process to

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request a step therapy exception determination. A group health plan may use its existing medical exceptions process in accordance with the provisions of Subsections D through I of this section to satisfy this requirement. The process shall be made easily accessible for enrollees and practitioners on the group health plan's publicly accessible website.

D. A group health plan shall expeditiously grant an exception to the group health plan's step therapy protocol, based on medical necessity and a clinically valid explanation from the patient's prescribing practitioner as to why a drug on the plan's formulary that is therapeutically equivalent to the prescribed drug should not be substituted for the prescribed drug, if:

(1) the prescription drug that is the subject of the exception request is contraindicated or will likely cause an adverse reaction by or physical or mental harm to the patient;

(2) the prescription drug that is the subject of the exception request is expected to be ineffective based on the known clinical characteristics of the patient and the known characteristics of the prescription drug regimen;

(3) while under the enrollee's current health coverage or previous health coverage, the enrollee has tried the prescription drug that is the subject of the exception

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request or another prescription drug in the same pharmacologic class or with the same mechanism of action as the prescription drug that is the subject of the exception request and that prescription drug was discontinued due to lack of efficacy or effectiveness, diminished effect or an adverse event; or

(4) the prescription drug required pursuant to the step therapy protocol is not in the best interest of the patient, based on clinical appropriateness, because the patient's use of the prescription drug is expected to:

(a) cause a significant barrier to the patient's adherence to or compliance with the patient's plan of care;

(b) worsen a comorbid condition of the patient; or

(c) decrease the patient's ability to achieve or maintain reasonable functional ability in performing daily activities.

E. Upon the granting of an exception to a group health plan's step therapy protocol, the group health plan administrator shall authorize continuing coverage STBTC → ~~for the life of the enrollee~~ ← STBTC for the prescription drug that is the subject of the exception request STBTC → for no less than the duration of the therapeutic effect of the drug ← STBTC . The group health plan shall include in its evidence of coverage

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language describing an enrollee's rights pursuant to this subsection.

F. A group health plan shall respond with its decision on an enrollee's exception request within seventy-two hours of receipt. In cases where exigent circumstances exist, a group health plan shall respond within twenty-four hours of receipt of the exception request. In the event the group health plan does not respond to an exception request within the time frames required pursuant to this subsection, the exception request shall be granted.

G. A group health plan administrator's denial of a request for an exception for step therapy protocols shall be subject to review and appeal pursuant to the Patient Protection Act.

H. After an enrollee has made an exception request in accordance with the provisions of this section, a group health plan shall authorize continued coverage of a prescription drug that is the subject of the exception request pending the determination of the exception request.

I. The provisions of this section shall not be construed to prevent a:

(1) group health plan from requiring a patient to try a STBTC→**biosimilar, interchangeable biologic or**←STBTC generic equivalent of a prescription drug before providing

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coverage for the equivalent brand-name prescription drug; or

(2) practitioner from prescribing a prescription drug that the practitioner has determined to be medically necessary.

~~[J. The provisions of this section shall apply only to a group health plan delivered, issued for delivery or renewed on or after January 1, 2019.~~

~~K.]~~ J. As used in this section, "medical necessity" or "medically necessary" means health care services determined by a practitioner, in consultation with the group health plan administrator, to be appropriate or necessary according to:

(1) any applicable, generally accepted principles and practices of good medical care;

(2) practice guidelines developed by the federal government or national or professional medical societies, boards or associations; or

(3) any applicable clinical protocols or practice guidelines developed by the group health plan consistent with federal, national and professional practice guidelines. These standards shall be applied to decisions related to the diagnosis or direct care and treatment of a physical or behavioral health condition, illness, injury or disease."

SECTION 2. Section 27-2-12.23 NMSA 1978 (being Laws 2018,

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Chapter 9, Section 2) is amended to read:

"27-2-12.23. MEDICAL ASSISTANCE--PRESCRIPTION DRUG
COVERAGE--STEP THERAPY PROTOCOLS--CLINICAL REVIEW CRITERIA--
EXCEPTIONS.--

A. By January 1, 2019, the secretary shall require any medical assistance plan for which any step therapy protocols are required to establish clinical review criteria for those step therapy protocols. The clinical review criteria shall be based on clinical practice guidelines that:

(1) recommend that the prescription drugs subject to step therapy protocols be taken in the specific sequence required by the step therapy protocol;

(2) are developed and endorsed by an interdisciplinary panel of experts that manages conflicts of interest among the members of the panel of experts by:

(a) requiring members to: 1) disclose any potential conflicts of interest with health care plans, medical assistance plans, health maintenance organizations, pharmaceutical manufacturers, pharmacy benefits managers and any other entities; and 2) recuse themselves if there is a conflict of interest; and

(b) using analytical and methodological experts to work to provide objectivity in data analysis and ranking of evidence through the preparation of evidence tables

and facilitating consensus;

(3) are based on high-quality studies, research and medical practice;

(4) are created pursuant to an explicit and transparent process that:

(a) minimizes bias and conflicts of interest;

(b) explains the relationship between treatment options and outcomes;

(c) rates the quality of the evidence supporting recommendations; and

(d) considers relevant patient subgroups and preferences; and

(5) take into account the needs of atypical patient populations and diagnoses.

B. In the absence of clinical guidelines that meet the requirements of Subsection A of this section, peer-reviewed publications may be substituted.

C. When a medical assistance plan restricts coverage of a prescription drug for the treatment of any medical condition through the use of a step therapy protocol, a recipient and the practitioner prescribing the prescription drug shall have access to a clear, readily accessible and convenient process to request a step therapy exception

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determination. A medical assistance plan may use its existing medical exceptions process in accordance with the provisions of Subsections D through I of this section to satisfy this requirement. The process shall be made easily accessible for recipients and practitioners on the medical assistance plan's publicly accessible website.

D. A medical assistance plan shall expeditiously grant an exception to the medical assistance plan's step therapy protocol, based on medical necessity and a clinically valid explanation from the patient's prescribing practitioner as to why a drug on the plan's formulary that is therapeutically equivalent to the prescribed drug should not be substituted for the prescribed drug, if:

(1) the prescription drug that is the subject of the exception request is contraindicated or will likely cause an adverse reaction by or physical or mental harm to the patient;

(2) the prescription drug that is the subject of the exception request is expected to be ineffective based on the known clinical characteristics of the patient and the known characteristics of the prescription drug regimen;

(3) while under the recipient's current medical assistance plan, or under the recipient's previous health coverage, the recipient has tried the prescription drug

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that is the subject of the exception request or another prescription drug in the same pharmacologic class or with the same mechanism of action as the prescription drug that is the subject of the exception request and that prescription drug was discontinued due to lack of efficacy or effectiveness, diminished effect or an adverse event; or

(4) the prescription drug required pursuant to the step therapy protocol is not in the best interest of the patient, based on clinical appropriateness, because the patient's use of the prescription drug is expected to:

(a) cause a significant barrier to the patient's adherence to or compliance with the patient's plan of care;

(b) worsen a comorbid condition of the patient; or

(c) decrease the patient's ability to achieve or maintain reasonable functional ability in performing daily activities.

E. Upon the granting of an exception to a medical assistance plan's step therapy protocol, a medical assistance plan shall authorize continuing coverage STBTC → ~~for the life of the patient~~ ← STBTC for the prescription drug that is the subject of the exception request STBTC → for no less than the duration of the therapeutic effect of the drug ← STBTC .

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F. A medical assistance plan shall respond with its decision on a recipient's exception request within seventy-two hours of receipt. In cases where exigent circumstances exist, a medical assistance plan shall respond within twenty-four hours of receipt of the exception request. In the event the medical assistance plan does not respond to an exception request within the time frames required pursuant to this subsection, the exception request shall be granted.

G. A medical assistance plan's denial of a request for an exception for step therapy protocols shall be subject to review and appeal pursuant to department rules.

H. After a recipient has made an exception request in accordance with the provisions of this section, a medical assistance plan shall authorize continued coverage of a prescription drug that is the subject of the exception request pending the determination of the exception request.

I. The provisions of this section shall not be construed to prevent:

(1) a medical assistance plan from requiring a patient to try a STBTC **→ biosimilar, interchangeable biologic** **or** ← STBTC generic equivalent of a prescription drug before providing coverage for the equivalent brand-name prescription drug; or

(2) a practitioner from prescribing a

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prescription drug that the practitioner has determined to be medically necessary.

J. As used in this section, "medical necessity" or "medically necessary" means health care services determined by a practitioner, in consultation with the medical assistance plan, to be appropriate or necessary, according to:

- (1) any applicable, generally accepted principles and practices of good medical care;
- (2) practice guidelines developed by the federal government or national or professional medical societies, boards or associations; or
- (3) any applicable clinical protocols or practice guidelines developed by the medical assistance plan consistent with federal, national and professional practice guidelines. These standards shall be applied to decisions related to the diagnosis or direct care and treatment of a physical or behavioral health condition, illness, injury or disease."

SECTION 3. Section 59A-22-53.1 NMSA 1978 (being Laws 2018, Chapter 9, Section 3) is amended to read:

"59A-22-53.1. PRESCRIPTION DRUG COVERAGE--STEP THERAPY PROTOCOLS--CLINICAL REVIEW CRITERIA--EXCEPTIONS.--

A. Each individual health insurance policy, health care plan and certificate of health insurance delivered or

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issued for delivery in this state that provides a prescription drug benefit for which any step therapy protocols are required shall establish clinical review criteria for those step therapy protocols. The clinical review criteria shall be based on clinical practice guidelines that:

(1) recommend that the prescription drugs subject to step therapy protocols be taken in the specific sequence required by the step therapy protocol;

(2) are developed and endorsed by an interdisciplinary panel of experts that manages conflicts of interest among the members of the panel of experts by:

(a) requiring members to: 1) disclose any potential conflicts of interest with insurers, health maintenance organizations, health care plans, pharmacy benefits managers and any other entities; and 2) recuse themselves if there is a conflict of interest; and

(b) using analytical and methodological experts to work to provide objectivity in data analysis and ranking of evidence through the preparation of evidence tables and facilitating consensus;

(3) are based on high-quality studies, research and medical practice;

(4) are created pursuant to an explicit and transparent process that:

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- (a) minimizes bias and conflicts of interest;
- (b) explains the relationship between treatment options and outcomes;
- (c) rates the quality of the evidence supporting recommendations; and
- (d) considers relevant patient subgroups and preferences; and
- (5) take into account the needs of atypical patient populations and diagnoses.

B. In the absence of clinical guidelines that meet the requirements of Subsection A of this section, peer-reviewed publications may be substituted.

C. When a health insurance policy, health care plan or certificate of insurance restricts coverage of a prescription drug for the treatment of any medical condition through the use of a step therapy protocol, an insured and the practitioner prescribing the prescription drug shall have access to a clear, readily accessible and convenient process to request a step therapy exception determination. An insurer may use its existing medical exceptions process in accordance with the provisions of Subsections D through I of this section to satisfy this requirement. The process shall be made easily accessible for insureds and practitioners on the insurer's

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publicly accessible website.

D. An insurer shall expeditiously grant an exception to the health insurance policy's, health care plan's or certificate of insurance's step therapy protocol, based on medical necessity and a clinically valid explanation from the patient's prescribing practitioner as to why a drug on the health insurance policy's, health care plan's or certificate of insurance's formulary that is therapeutically equivalent to the prescribed drug should not be substituted for the prescribed drug, if:

(1) the prescription drug that is the subject of the exception request is contraindicated or will likely cause an adverse reaction by or physical or mental harm to the patient;

(2) the prescription drug that is the subject of the exception request is expected to be ineffective based on the known clinical characteristics of the patient and the known characteristics of the prescription drug regimen;

(3) while under the insured's current health insurance policy, health care plan or certificate of insurance, or under the insured's previous health coverage, the insured has tried the prescription drug that is the subject of the exception request or another prescription drug in the same pharmacologic class or with the same mechanism of action as the

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prescription drug that is the subject of the exception request and that prescription drug was discontinued due to lack of efficacy or effectiveness, diminished effect or an adverse event; or

(4) the prescription drug required pursuant to the step therapy protocol is not in the best interest of the patient, based on clinical appropriateness, because the patient's use of the prescription drug is expected to:

(a) cause a significant barrier to the patient's adherence to or compliance with the patient's plan of care;

(b) worsen a comorbid condition of the patient; or

(c) decrease the patient's ability to achieve or maintain reasonable functional ability in performing daily activities.

E. Upon the granting of an exception to a health insurance policy's, health care plan's or certificate of insurance's step therapy protocol, an insurer shall authorize coverage ~~STBTC~~ **for the life of the insured** ~~STBTC~~ for the prescription drug that is the subject of the exception request ~~STBTC~~ **for no less than the duration of the therapeutic effect of the drug** ~~STBTC~~ . An insurer shall include in its evidence of coverage language describing an insured's rights pursuant to

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this subsection.

F. An insurer shall respond with its decision on an insured's exception request within seventy-two hours of receipt. In cases where exigent circumstances exist, an insurer shall respond within twenty-four hours of receipt of the exception request. In the event the insurer does not respond to an exception request within the time frames required pursuant to this subsection, the exception request shall be granted.

G. An insurer's denial of a request for an exception for step therapy protocols shall be subject to review and appeal pursuant to the Patient Protection Act.

H. After an insured has made an exception request in accordance with the provisions of this section, an insurer shall authorize continued coverage of a prescription drug that is the subject of the exception request pending the determination of the exception request.

I. The provisions of this section shall not be construed to prevent:

(1) a health insurance policy, health care plan or certificate of insurance from requiring a patient to try a STBTC → biosimilar, interchangeable biologic or ← STBTC generic equivalent of a prescription drug before providing coverage for the equivalent brand-name prescription drug; or

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(2) a practitioner from prescribing a prescription drug that the practitioner has determined to be medically necessary.

~~[J. The provisions of this section shall apply only to a health insurance policy, health care plan or certificate of insurance delivered, issued for delivery or renewed on or after January 1, 2019.]~~

~~K.]~~ J. The superintendent shall promulgate rules as may be necessary to appropriately implement the provisions of this section.

~~[L.]~~ K. Nothing in this section shall be interpreted to interfere with the superintendent's authority to regulate prescription drug coverage benefits under other state and federal law.

~~[M.]~~ L. As used in this section, "medical necessity" or "medically necessary" means health care services determined by a practitioner, in consultation with the insurer, to be appropriate or necessary, according to:

(1) any applicable, generally accepted principles and practices of good medical care;

(2) practice guidelines developed by the federal government or national or professional medical societies, boards or associations; or

(3) any applicable clinical protocols or

practice guidelines developed by the insurer consistent with federal, national and professional practice guidelines. These standards shall be applied to decisions related to the diagnosis or direct care and treatment of a physical or behavioral health condition, illness, injury or disease."

SECTION 4. Section 59A-22B-8 NMSA 1978 (being Laws 2023, Chapter 114, Section 13) is amended to read:

"59A-22B-8. PRIOR AUTHORIZATION FOR PRESCRIPTION DRUGS OR STEP THERAPY FOR ~~[SUBSTANCE USE DISORDER]~~ CERTAIN CONDITIONS PROHIBITED.--

A. Coverage for medication approved by the federal food and drug administration that is prescribed for the treatment of an autoimmune disorder, STBTC→~~a behavioral health condition,~~←STBTC cancer or a substance use disorder, pursuant to a medical necessity determination, shall not be subject to prior authorization, except in cases in which a STBTC→biosimilar, interchangeable biologic or←STBTC generic version is available.

B. A health insurer shall not impose step therapy requirements before authorizing coverage for medication approved by the federal food and drug administration that is prescribed for the treatment of an autoimmune disorder, STBTC→~~a behavioral health condition,~~←STBTC cancer or a substance use disorder, pursuant to a medical necessity determination, except

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in cases in which a STBTC→**biosimilar, interchangeable biologic**
or←STBTC generic version is available."

SECTION 5. Section 59A-46-52.1 NMSA 1978 (being Laws
2018, Chapter 9, Section 5) is amended to read:

"59A-46-52.1. PRESCRIPTION DRUG COVERAGE--STEP THERAPY
PROTOCOLS--CLINICAL REVIEW CRITERIA--EXCEPTIONS.--

A. Each individual or group health maintenance
organization contract delivered or issued for delivery in this
state that provides a prescription drug benefit for which any
step therapy protocols are required shall establish clinical
review criteria for those step therapy protocols. The clinical
review criteria shall be based on clinical practice guidelines
that:

(1) recommend that the prescription drugs
subject to step therapy protocols be taken in the specific
sequence required by the step therapy protocol;

(2) are developed and endorsed by an
interdisciplinary panel of experts that manages conflicts of
interest among the members of the panel of experts by:

(a) requiring members to: 1) disclose
any potential conflicts of interest with carriers, insurers,
health care plans, pharmaceutical manufacturers, pharmacy
benefits managers and any other entities; and 2) recuse
themselves if there is a conflict of interest; and

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(b) using analytical and methodological experts to work to provide objectivity in data analysis and ranking of evidence through the preparation of evidence tables and facilitating consensus;

(3) are based on high-quality studies, research and medical practice;

(4) are created pursuant to an explicit and transparent process that:

(a) minimizes bias and conflicts of interest;

(b) explains the relationship between treatment options and outcomes;

(c) rates the quality of the evidence supporting recommendations; and

(d) considers relevant patient subgroups and preferences; and

(5) take into account the needs of atypical patient populations and diagnoses.

B. In the absence of clinical guidelines that meet the requirements of Subsection A of this section, peer-reviewed publications may be substituted.

C. When a health maintenance organization contract restricts coverage of a prescription drug for the treatment of any medical condition through the use of a step therapy

protocol, an enrollee and the practitioner prescribing the prescription drug shall have access to a clear, readily accessible and convenient process to request a step therapy exception determination. A carrier may use its existing medical exceptions process in accordance with the provisions of Subsections D through I of this section to satisfy this requirement. The process shall be made easily accessible for enrollees and practitioners on the carrier's publicly accessible website.

D. A carrier shall expeditiously grant an exception to the health maintenance organization contract's step therapy protocol, based on medical necessity and a clinically valid explanation from the patient's prescribing practitioner as to why a drug on the health maintenance organization contract's formulary that is therapeutically equivalent to the prescribed drug should not be substituted for the prescribed drug, if:

(1) the prescription drug that is the subject of the exception request is contraindicated or will likely cause an adverse reaction by or physical or mental harm to the patient;

(2) the prescription drug that is the subject of the exception request is expected to be ineffective based on the known clinical characteristics of the patient and the known characteristics of the prescription drug regimen;

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(3) while under the enrollee's current health maintenance organization contract, or under the enrollee's previous health coverage, the enrollee has tried the prescription drug that is the subject of the exception request or another prescription drug in the same pharmacologic class or with the same mechanism of action as the prescription drug that is the subject of the exception request and that prescription drug was discontinued due to lack of efficacy or effectiveness, diminished effect or an adverse event; or

(4) the prescription drug required pursuant to the step therapy protocol is not in the best interest of the patient, based on clinical appropriateness, because the patient's use of the prescription drug is expected to:

(a) cause a significant barrier to the patient's adherence to or compliance with the patient's plan of care;

(b) worsen a comorbid condition of the patient; or

(c) decrease the patient's ability to achieve or maintain reasonable functional ability in performing daily activities.

E. Upon the granting of an exception to a health maintenance organization contract's step therapy protocol, a carrier shall authorize coverage STBTC → ~~for the lifetime of the~~

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~~enrollee~~ ← STBTC for the prescription drug that is the subject of the exception request STBTC → **for no less than the duration of the therapeutic effect of the drug** ← STBTC . A carrier shall include in its evidence of coverage language describing an enrollee's rights pursuant to this subsection.

F. A carrier shall respond with its decision on an enrollee's exception request within seventy-two hours of receipt. In cases where exigent circumstances exist, a carrier shall respond within twenty-four hours of receipt of the exception request. In the event the carrier does not respond to an exception request within the time frames required pursuant to this subsection, the exception request shall be granted.

G. A carrier's denial of a request for an exception for step therapy protocols shall be subject to review and appeal pursuant to the Patient Protection Act.

H. After an enrollee has made an exception request in accordance with the provisions of this section, a carrier shall authorize continued coverage of a prescription drug that is the subject of the exception request pending the determination of the exception request.

I. The provisions of this section shall not be construed to prevent:

- (1) a health maintenance organization contract

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from requiring a patient to try a ~~STBTC~~→**biosimilar**,
interchangeable biologic or←~~STBTC~~ generic equivalent of a
prescription drug before providing coverage for the equivalent
brand-name prescription drug; or

(2) a practitioner from prescribing a
prescription drug that the practitioner has determined to be
medically necessary.

~~[J. The provisions of this section shall apply only
to a health maintenance organization contract delivered, issued
for delivery or renewed on or after January 1, 2019.~~

~~K.]~~ J. The superintendent shall promulgate rules as
may be necessary to appropriately implement the provisions of
this section.

~~[L.]~~ K. Nothing in this section shall be
interpreted to interfere with the superintendent's authority to
regulate prescription drug coverage benefits under other state
and federal law.

~~[M.]~~ L. As used in this section, "medical
necessity" or "medically necessary" means health care services
determined by a practitioner, in consultation with the carrier,
to be appropriate or necessary, according to:

(1) any applicable, generally accepted
principles and practices of good medical care;

(2) practice guidelines developed by the

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federal government or national or professional medical societies, boards or associations; or

(3) any applicable clinical protocols or practice guidelines developed by the carrier consistent with federal, national and professional practice guidelines. These standards shall be applied to decisions related to the diagnosis or direct care and treatment of a physical or behavioral health condition, illness, injury or disease."

SECTION 6. Section 59A-47-47.1 NMSA 1978 (being Laws 2018, Chapter 9, Section 6) is amended to read:

"59A-47-47.1. PRESCRIPTION DRUG COVERAGE--STEP THERAPY PROTOCOLS--CLINICAL REVIEW CRITERIA--EXCEPTIONS.--

A. Each individual or group nonprofit health care plan contract delivered or issued for delivery in this state that provides a prescription drug benefit for which any step therapy protocols are required shall establish clinical review criteria for those step therapy protocols. The clinical review criteria shall be based on clinical practice guidelines that:

(1) recommend that the prescription drugs subject to step therapy protocols be taken in the specific sequence required by the step therapy protocol;

(2) are developed and endorsed by an interdisciplinary panel of experts that manages conflicts of interest among the members of the panel of experts by:

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(a) requiring members to: 1) disclose any potential conflicts of interest with health care plans, insurers, health maintenance organizations, pharmaceutical manufacturers, pharmacy benefits managers and any other entities; and 2) recuse themselves if there is a conflict of interest; and

(b) using analytical and methodological experts to work to provide objectivity in data analysis and ranking of evidence through the preparation of evidence tables and facilitating consensus;

(3) are based on high-quality studies, research and medical practice;

(4) are created pursuant to an explicit and transparent process that:

(a) minimizes bias and conflicts of interest;

(b) explains the relationship between treatment options and outcomes;

(c) rates the quality of the evidence supporting recommendations; and

(d) considers relevant patient subgroups and preferences; and

(5) take into account the needs of atypical patient populations and diagnoses.

B. In the absence of clinical guidelines that meet the requirements of Subsection A of this section, peer-reviewed publications may be substituted.

C. When a health care plan restricts coverage of a prescription drug for the treatment of any medical condition through the use of a step therapy protocol, a subscriber and the practitioner prescribing the prescription drug shall have access to a clear, readily accessible and convenient process to request a step therapy exception determination. A health care plan may use its existing medical exceptions process in accordance with the provisions of Subsections D through I of this section to satisfy this requirement. The process shall be made easily accessible for subscribers and practitioners on the health care plan's publicly accessible website.

D. A health care plan shall expeditiously grant an exception to the health care plan's step therapy protocol, based on medical necessity and a clinically valid explanation from the patient's prescribing practitioner as to why a drug on the health care plan's formulary that is therapeutically equivalent to the prescribed drug should not be substituted for the prescribed drug, if:

(1) the prescription drug that is the subject of the exception request is contraindicated or will likely cause an adverse reaction by or physical or mental harm to the

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patient;

(2) the prescription drug that is the subject of the exception request is expected to be ineffective based on the known clinical characteristics of the patient and the known characteristics of the prescription drug regimen;

(3) while under the subscriber's current health care plan, or under the subscriber's previous health coverage, the subscriber has tried the prescription drug that is the subject of the exception request or another prescription drug in the same pharmacologic class or with the same mechanism of action as the prescription drug that is the subject of the exception request and that prescription drug was discontinued due to lack of efficacy or effectiveness, diminished effect or an adverse event; or

(4) the prescription drug required pursuant to the step therapy protocol is not in the best interest of the patient, based on clinical appropriateness, because the patient's use of the prescription drug is expected to:

(a) cause a significant barrier to the patient's adherence to or compliance with the patient's plan of care;

(b) worsen a comorbid condition of the patient; or

(c) decrease the patient's ability to

achieve or maintain reasonable functional ability in performing daily activities.

E. Upon the granting of an exception to a health care plan's step therapy protocol, a health care plan shall authorize coverage ~~STBTC~~ for the lifetime of the subscriber ~~STBTC~~ for the prescription drug that is the subject of the exception request ~~STBTC~~ for no less than the duration of the therapeutic effect of the drug ~~STBTC~~ . A health care plan shall include in its evidence of coverage language describing a subscriber's rights pursuant to this subsection.

F. A health care plan shall respond with its decision on a subscriber's exception request within seventy-two hours of receipt. In cases where exigent circumstances exist, a health care plan shall respond within twenty-four hours of receipt of the exception request. In the event the insurer does not respond to an exception request within the time frames required pursuant to this subsection, the exception request shall be granted.

G. A health care plan's denial of a request for an exception for step therapy protocols shall be subject to review and appeal pursuant to the Patient Protection Act.

H. After a subscriber has made an exception request in accordance with the provisions of this section, a health care plan shall authorize continued coverage of a prescription

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drug that is the subject of the exception request pending the determination of the exception request.

I. The provisions of this section shall not be construed to prevent:

(1) a health care plan from requiring a patient to try a ~~STBTC~~ **biosimilar, interchangeable biologic** ~~or~~ ~~STBTC~~ generic equivalent of a prescription drug before providing coverage for the equivalent brand-name prescription drug; or

(2) a practitioner from prescribing a prescription drug that the practitioner has determined to be medically necessary.

~~[J. The provisions of this section shall apply only to a health care plan delivered, issued for delivery or renewed on or after January 1, 2019.~~

~~K.]~~ J. The superintendent shall promulgate rules as may be necessary to appropriately implement the provisions of this section.

~~[L.]~~ K. Nothing in this section shall be interpreted to interfere with the superintendent's authority to regulate prescription drug coverage benefits under other state and federal law.

~~[M.]~~ L. As used in this section, "medical necessity" or "medically necessary" means health care services

determined by a practitioner, in consultation with the health care plan, to be appropriate or necessary, according to:

- (1) any applicable, generally accepted principles and practices of good medical care;
- (2) practice guidelines developed by the federal government or national or professional medical societies, boards or associations; or
- (3) any applicable clinical protocols or practice guidelines developed by the health care plan consistent with federal, national and professional practice guidelines. These standards shall be applied to decisions related to the diagnosis or direct care and treatment of a physical or behavioral health condition, illness, injury or disease."

SECTION 7. EXCEPTIONS.--The provisions of Sections 1 and 3 through 6 of this act do not apply to short-term plans subject to the Short-Term Health Plan and Excepted Benefit Act.

SECTION 8. APPLICABILITY.--The provisions of this act apply to group health insurance policies, health care plans or certificates of health insurance, other than small group health plans, that are delivered, issued for delivery or renewed in this state on or after January 1, 2025.

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