As Introduced

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Representative Miller, K.
Cosponsors: Representatives Rogers, Manning, Johnson, Cutrona, Hillyer, Troy, Galonski, Somani, Denson, Manchester, Dell'Aquila, Dean, Cross

A BILL

To amend section 5160.34 and to enact sections 1751.721, 1751.722, 1751.723, 3923.042, 3923.043, 3923.044, 5160.341, and 5160.342 of the Revised Code to establish an exemption to prior authorization requirements.

BE IT ENACTED BY THE GENERAL ASSEMBLY OF THE STATE OF OHIO:

Section 1. That section 5160.34 be amended and sections 1751.721, 1751.722, 1751.723, 3923.042, 3923.043, 3923.044, 5160.341, and 5160.342 of the Revised Code be enacted to read as follows:

Sec. 1751.721. (A) A health insuring corporation that applies a prior authorization requirement shall make prior authorization data available on its public web site in a readily accessible format.

(B) The data shall include all of the following information:

(1) The specialty of the health care provider requesting the prior authorization;
(2) Whether the prior authorization is for a health care service, a medical device, or a drug;

(3) The indication for use of the service, device, or drug under the prior authorization;

(4) If the prior authorization request was denied, the reason for the denial;

(5) If the approval or denial of a prior authorization request was appealed and the result of the appeal;

(6) The amount of time between the submission of a prior authorization request and the response from the corporation.

Sec. 1751.722. (A)(1) If a health insuring corporation has a prior authorization requirement for a health care service, medical device, or drug and, during the previous twelve-month period, the corporation approved at least eighty per cent of the prior authorization requests submitted by a health care provider for that service, device, or drug, the insurer or its designee shall not require the health care provider to comply with the requirement for that service, device, or drug.

(2) Such an exemption shall be provided for not less than twelve months.

(3) Nothing in this section shall be construed as prohibiting a corporation from establishing an exemption period of more than twelve months.

(B)(1) A health care provider that does not receive an exemption under division (A) of this section may request that the corporation provide evidence to the provider supporting its decision to not grant an exemption.

(2) The health care provider may make such a request at
any time, but it may make not more than one such request for the
same service, device, or drug in a calendar year.

(3) A health insuring corporation shall comply with such a
request.

(C) A health care provider may appeal a health insuring
corporation's decision to deny an exemption.

(D) A health insuring corporation shall not require a
health care provider to request an exemption provided under
division (A) of this section.

(E) When an exemption is granted under division (A) of
this section for a health care service, medical device, or drug,
the corporation shall notify the health care provider in
question. The notice shall be in writing and include all of the
following information:

(1) A statement that the health care provider qualifies
for an exemption to a prior authorization requirement;

(2) The health care service, medical device, or drug to
which the exemption applies;

(3) The dates the exemption will begin and end.

(F)(1) At the end of the twelve-month exemption period, a
health insuring corporation may evaluate an exemption it has
granted under division (A) of this section.

(2)(a) A corporation conducting such an evaluation shall
review ten claims submitted to the corporation, selected at
random, for the health care service, medical device, or drug in
question.

(b) The reviewed claims shall be from the immediately
preceding three months. If there are not ten relevant claims in
the preceding three months, the corporation may review earlier
claims.

(3)(a) If less than eighty per cent of the claims reviewed
would have been approved based on medical necessity, then the
corporation may revoke the exemption provided under division (A)
of this section.

(b) A corporation that is revoking an exemption shall
provide the health care provider with both of the following:

(i) The information it relied upon in making its
determination;

(ii) A plain language explanation of how to appeal the
decision.

(4) A corporation shall not evaluate a health care
provider's exemption relating to a particular service, device,
or drug more than once every twelve months.

(5) Nothing in this section shall be construed as
requiring a corporation to evaluate an existing exemption.

(G) If an exemption is revoked and not appealed, the
exemption shall remain in effect until thirty days after the
date the corporation notifies the health care provider of the
corporation's decision to revoke the exemption.

(H) A health care provider may appeal the revocation of an
exemption within thirty days of receiving notice of the
revocation. If the health care provider appeals the revocation
and the revocation is upheld, the exemption remains in effect
until five days after the date the revocation is upheld.

(I) A decision to revoke or deny an exemption shall only
be made by a health care provider licensed in this state who
practices the same or a similar specialty as the health care
provider being considered for an exemption and who has
experience in providing the service, device, or drug to which
the exemption or potential exemption applies.

(J) Nothing in this section shall be construed as
prohibiting a health insuring corporation from making an
administrative denial of a claim.

Sec. 1751.723. (A) A series of violations of section
1751.721 or 1751.722 of the Revised Code that, taken together,
constitute a practice or pattern shall be considered an unfair
and deceptive practice under sections 3901.19 to 3901.26 of the
Revised Code.

(B) Notwithstanding division (F) of section 121.95 of the
Revised Code, the superintendent of insurance may adopt rules as
necessary to carry out the requirements of sections 1751.721 to
1751.723 of the Revised Code.

Sec. 3923.042. (A) A sickness and accident insurer that
applies a prior authorization requirement shall make prior
authorization data available on its public web site in a readily
accessible format.

(B) The data shall include all of the following
information:

(1) The specialty of the health care provider requesting
the prior authorization;

(2) Whether the prior authorization is for a medical
service, a medical device, or a drug;

(3) The indication for use of the service, device, or drug
under the prior authorization;

(4) If the prior authorization request was denied, the reason for the denial;

(5) If the approval or denial of a prior authorization request was appealed and the result of the appeal;

(6) The amount of time between the submission of a prior authorization request and the response from the insurer.

Sec. 3923.043. (A)(1) If a sickness and accident insurer has a prior authorization requirement for a health care service, medical device, or drug and, during the previous twelve-month period, the insurer approved at least eighty per cent of the prior authorization requests submitted by a health care provider for that service, device, or drug, the insurer or its designee shall not require the health care provider to comply with the requirement for that service, device, or drug.

(2) Such an exemption shall be provided for not less than twelve months.

(3) Nothing in this section shall be construed as prohibiting an insurer from establishing an exemption period of more than twelve months.

(B)(1) A health care provider that does not receive an exemption under division (A) of this section may request that the sickness and accident insurer provide evidence to the provider supporting its decision to not grant an exemption.

(2) The insurer may make the request at any time, but it may make not more than one such request for the same service, device, or drug in a calendar year.

(3) A sickness and accident insurer shall comply with such
a request.

(C) A health care provider may appeal a sickness and accident insurer's decision to deny an exemption.

(D) A sickness and accident insurer shall not require a health care provider to request an exemption provided under division (A) of this section.

(E) When an exemption is granted under division (A) of this section for a health care service, medical device, or drug, the sickness and accident insurer shall notify the health care provider in question. The notice shall be in writing and include all of the following information:

(1) A statement that the health care provider qualifies for an exemption to a prior authorization requirement;

(2) The health care service, medical device, or drug to which the exemption applies;

(3) The dates the exemption will begin and end.

(F)(1) At the end of the twelve-month exemption period, a sickness and accident insurer may evaluate an exemption it has granted under division (A) of this section.

(2)(a) An insurer conducting such an evaluation shall review ten claims submitted to the insurer, selected at random, for the health care service, medical device, or drug in question.

(b) The reviewed claims shall be from the immediately preceding three months. If there are not ten relevant claims in the preceding three months, the insurer may review earlier claims.
(3)(a) If less than eighty per cent of the claims reviewed would have been approved based on medical necessity, then the insurer may revoke the exemption provided under division (A) of this section.

(b) An insurer that is revoking an exemption shall provide the health care provider with both of the following:

(i) The information it relied upon in making its determination;

(ii) A plain language explanation of how to appeal the decision.

(4) An insurer shall not evaluate a health care provider's exemption relating to a particular service, device, or drug more than once every twelve months.

(5) Nothing in this section shall be construed as requiring an insurer to evaluate an existing exemption.

(G) If an exemption is revoked and not appealed, the exemption shall remain in effect until thirty days after the date the sickness and accident insurer notifies the health care provider of the insurer's decision to revoke the exemption.

(H) A health care provider may appeal the revocation of an exemption within thirty days after receiving notification of the revocation. If a health care provider appeals a revocation and the revocation is upheld, the exemption remains in effect until five days after the date the revocation is upheld.

(I) A decision to revoke or deny an exemption shall only be made by a health care provider licensed in this state who practices the same or a similar specialty as the health care provider being considered for an exemption and who has
experience in providing the service, device, or drug to which the exemption or potential exemption applies.

(J) Nothing in this section shall be construed as prohibiting a sickness and accident insurer from making an administrative denial of a claim.

**Sec. 3923.044.** (A) A series of violations of section 3923.042 or 3923.043 of the Revised Code that, taken together, constitute a practice or pattern shall be considered an unfair and deceptive practice under sections 3901.19 to 3901.26 of the Revised Code.

(B) Notwithstanding division (F) of section 121.95 of the Revised Code, the superintendent of insurance may adopt rules as necessary to carry out the requirements of sections 3923.042 to 3923.044 of the Revised Code.

**Sec. 5160.34.** (A) As used in this section sections 5160.34 to 5160.342 of the Revised Code:

(1) "Chronic condition" means a medical condition that has persisted after reasonable efforts have been made to relieve or cure its cause and has continued, either continuously or episodically, for longer than six continuous months.

(2) "Clinical peer" means a health care provider in the same, or in a similar, specialty that typically manages the medical condition, procedure, or treatment under review.

(3) "Emergency services" has the same meaning as in section 1753.28 of the Revised Code.

(4) "Prior authorization requirement" means any practice implemented by a medical assistance program in which coverage of a health care service, device, or drug is dependent upon a
medical assistance recipient or a health care provider, receiving approval from the department of medicaid or its designee, including a medicaid managed care organization, prior to the service, device, or drug being performed, received, or prescribed, as applicable. "Prior authorization" includes prospective or utilization review procedures conducted prior to providing a health care service, medical device, or drug.

(5) "Urgent care services" means a medical care or other service for a condition where application of the timeframe for making routine or non-life threatening care determinations is either of the following:

(a) Could seriously jeopardize the life, health, or safety of the recipient or others due to the recipient's psychological state;

(b) In the opinion of a practitioner with knowledge of the recipient's medical or behavioral condition, would subject the recipient to adverse health consequences without the care or treatment that is the subject of the request.

(6) "Utilization review" and "utilization review organization" have the same meaning as in section 1751.77 of the Revised Code.

(B) If a medical assistance program has a prior authorization requirement, the department of medicaid or its designee, including a medicaid managed care organization, shall do all of the following:

(1) On or before January 1, 2018, permit a health care provider to access the prior authorization form through the applicable electronic software system.

(2)(a) On or before January 1, 2018, permit the department
or its designee to accept and respond to prior prescription benefit authorization requests through a secure electronic transmission.

(b) On or before January 1, 2018, the department or its designee shall accept and respond to prior prescription benefit authorization requests through a secure electronic transmission using NCPDP SCRIPT standard ePA transactions, and for prior medical benefit authorization requests through a secure electronic transmission using standards established by the council for affordable quality health care on operating rules for information exchange or its successor.

(c) For purposes of division (B)(2) of this section, neither of the following shall be considered a secure electronic transmission:

(i) A facsimile;

(ii) A proprietary payer portal for prescription drug requests that does not use NCPDP SCRIPT standard.

(3) On or before January 1, 2018, a health care provider and the department of medicaid or its designee may enter into a contractual arrangement under which the department or its designee agrees to process prior authorization requests that are not submitted electronically because of the financial hardship that electronic submission of prior authorization requests would create for the provider or if internet connectivity is limited or unavailable where the provider is located.

(4)(a) On or before January 1, 2018, if the health care provider submits the request for prior authorization electronically as described in divisions (B)(1) and (2) of this section, the department or its designee shall respond to all
prior authorization requests within forty-eight hours for urgent care services, or ten calendar days for any prior authorization request that is not for an urgent care service, of the time the request is received by the department or its designee. Division (B)(4) of this section does not apply to emergency services.

(b) The response required under division (B)(4)(a) of this section shall indicate whether the request is approved or denied. If the prior authorization is denied, the department or its designee shall provide the specific reason for the denial.

(c) If the prior authorization request is incomplete, the department or its designee shall indicate the specific additional information that is required to process the request.

(5)(a) On or before January 1, 2018, if a health care provider submits a prior authorization request as described in divisions (B)(1) and (2) of this section, the department or its designee shall provide an electronic receipt to the health care provider acknowledging that the prior authorization request was received.

(b) On or before January 1, 2018, if the department or its designee requests additional information that is required to process a prior authorization request as described in division (B)(4)(c) of this section, the health care provider shall provide an electronic receipt to the department or its designee acknowledging that the request for additional information was received.

(6)(a) On or before January 1, 2017, honor a prior authorization approval for an approved drug for the lesser of the following from the date of approval:

(i) Twelve months;
(ii) The last day of the medical assistance recipient's eligibility for the medical assistance program.

(b) The duration of all other prior authorization approvals shall be dictated by the medical assistance program.

(c) The department or its designee, in relation to prior approval under division (B)(6)(a) of this section, may require a health care provider to submit information to the department or its designee indicating that the patient's chronic condition has not changed.

(i) The request for information by the department or its designee and the response by the health care provider shall be in an electronic format, which may be by electronic mail or other electronic communication.

(ii) The frequency of the submission of requested information shall be consistent with medical or scientific evidence as defined in section 3922.01 of the Revised Code, but shall not be required more frequently than quarterly.

(iii) If the health care provider does not respond within five calendar days from the date the request was received, the insurer or plan may terminate the twelve-month approval.

(d) A twelve-month approval provided under division (B)(6)(a) of this section is no longer valid and automatically terminates if there are changes to federal or state laws or federal regulatory guidance or compliance information prescribing that the drug in question is no longer approved or safe for the intended purpose.

(e) A twelve-month approval provided under division (B)(6)(a) of this section does not apply to and is not required for any of the following:
(i) Medications that are prescribed for a non-maintenance condition;

(ii) Medications that have a typical treatment of less than one year;

(iii) Medications that require an initial trial period to determine effectiveness and tolerability, beyond which a one-year, or greater, prior authorization period will be given;

(iv) Medications where there is medical or scientific evidence as defined in section 3922.01 of the Revised Code that do not support a twelve-month prior approval;

(v) Medications that are a schedule I or II controlled substance or any opioid analgesic or benzodiazepine, as defined in section 3719.01 of the Revised Code;

(vi) Medications that are not prescribed by an in-network provider as part of a care management program.

(?) On or before January 1, 2017, the department or its designee may, but is not required to, provide the twelve-month approval prescribed in division (B)(6)(a) of this section for a prescription drug that meets either of the following:

(a) The drug is prescribed or administered to treat a rare medical condition and pursuant to medical or scientific evidence as defined in section 3922.01 of the Revised Code.

(b) Medications that are controlled substances not included in division (B)(6)(e)(v) of this section.

For purposes of division (B)(?) of this section, "rare medical condition" means any disease or condition that affects fewer than two-hundred thousand individuals in the United States.
(8) Nothing in division (B)(6) or (7) of this section prohibits the substitution, in accordance with section 4729.38 of the Revised Code, of any drug that has received a twelve-month approval under division (B)(6)(a) of this section when there is a release of either of the following:

(a) A United States food and drug administration approved comparable brand product or a generic counterpart of a brand product that is listed as therapeutically equivalent in the United States food and drug administration's publication titled approved drug products with therapeutic equivalence evaluations;

(b) An interchangeable biological product, as defined in section 3715.01 of the Revised Code.

(9)(a) On or after January 1, 2017, upon written request, the department or its designee shall permit a retrospective review for a claim that is submitted for a service where prior authorization was required, but not obtained if the service in question meets all of the following:

(i) The service is directly related to another service for which prior approval has already been obtained and that has already been performed.

(ii) The new service was not known to be needed at the time the original prior authorized service was performed.

(iii) The need for the new service was revealed at the time the original authorized service was performed.

(b) Once the written request and all necessary information is received, the department or its designee shall review the claim for coverage and medical necessity. The department or its designee shall not deny a claim for such a new service based solely on the fact that a prior authorization approval was not
received for the new service in question.

(10)(a) On or before January 1, 2017, disclose to all participating health care providers any new prior authorization requirement at least thirty days prior to the effective date of the new requirement.

(b) The notice may be sent via electronic mail or standard mail and shall be conspicuously entitled "Notice of Changes to Prior Authorization Requirements." The notice is not required to contain a complete listing of all changes made to the prior authorization requirements, but shall include specific information on where the health care provider may locate the information on the department's or its designee's web site or, if applicable, the department's or its designee's portal.

(c) All participating health care providers shall promptly notify the department or its designee of any changes to the health care provider's electronic mail or standard mail address.

(11)(a) On or before January 1, 2017, make available to all participating health care providers on its web site or provider portal a listing of its prior authorization requirements, including specific information or documentation that a provider must submit in order for the prior authorization request to be considered complete.

(b) Make available on its web site information about the medical assistance programs offered in this state that clearly identifies specific services, drugs, or devices to which a prior authorization requirement exists.

(12) On or before January 1, 2018, establish a streamlined appeal process relating to adverse prior authorization determinations that shall include all of the following:
(a) For urgent care services, the appeal shall be considered within forty-eight hours after the department or its designee receives the appeal.

(b) For all other matters, the appeal shall be considered within ten calendar days after the department or its designee receives the appeal.

(c) The appeal shall be between the health care provider requesting the service in question and a clinical peer appointed by or contracted by the department or the department's designee.

(d) If the appeal does not resolve the disagreement, the appeal procedures shall permit the recipient to further appeal in accordance with section 5160.31 of the Revised Code.

(C) Beginning January 1, 2017, except in cases of fraudulent or materially incorrect information, the department or its designee shall not retroactively deny a prior authorization for a health care service, drug, or device when all of the following are met:

(1) The health care provider submits a prior authorization request to the department or its designee for a health care service, drug, or device.

(2) The department or its designee approves the prior authorization request after determining that all of the following are true:

(a) The recipient is eligible for the health care service, drug, or device under the medical assistance program.

(b) The health care service, drug, or device is covered by the medical assistance program.

(c) The health care service, drug, or device meets the
department's standards for medical necessity and prior
authorization.

(3) The health care provider renders the health care
service, drug, or device pursuant to the approved prior
authorization request and all of the terms and conditions of the
health care provider's contract with the department or the
department's designee.

(4) On the date the health care provider renders the prior
approved health care service, drug, or device, all of the
following are true:

(a) The recipient is eligible for the medical assistance
program.

(b) The recipient's condition or circumstances related to
the recipient's care has not changed.

(c) The health care provider submits an accurate claim
that matches the information submitted by the health care
provider in the approved prior authorization request.

(5) If the health care provider submits a claim that
includes an unintentional error and the error results in a claim
that does not match the information originally submitted by the
health care provider in the approved prior authorization
request, upon receiving a denial of services from the department
or its designee, the health care provider may resubmit the claim
pursuant to division (C) of this section with the information
that matches the information included in the approved prior
authorization.

(D) Any provision of a contractual arrangement entered
into between the department or its designee and a health care
provider or recipient that is contrary to divisions (A) to (C)
of this section is unenforceable.

(E) The director of medicaid may adopt rules in accordance with Chapter 119. of the Revised Code as necessary to implement the provisions of this section and section 5160.342 of the Revised Code.

Sec. 5160.341. (A) If the department or its designee applies a prior authorization requirement, it shall make prior authorization data available on its public web site in a readily accessible format.

(B) The data shall include all of the following information:

(1) The specialty of the health care provider requesting the prior authorization;

(2) Whether the prior authorization is for a health care service, a medical device, or a drug;

(3) The indication for use of the service, device, or drug under the prior authorization;

(4) If the prior authorization request was denied, the reason for the denial;

(5) If the approval or denial of a prior authorization request was appealed and the result of the appeal;

(6) The amount of time between the submission of a prior authorization request and the response from the department or its designee.

Sec. 5160.342. (A)(1) If a medical assistance program has a prior authorization requirement for a health care service, medical device, or drug and, during the previous twelve-month
period, the department of medicaid or its designee approved at
least eighty per cent of the prior authorization requests
submitted by a health care provider for that service, device, or
drug, the department or its designee shall not require the
health care provider to comply with the requirement for that
service, device, or drug.

(2) Such an exemption shall be provided for not less than
twelve months.

(3) Nothing in this section shall be construed as
prohibiting the department or its designee from establishing an
exemption period of more than twelve months.

(B)(1) A health care provider that does not receive an
exemption under division (A) of this section may request that
the department or the department's designee provide evidence to
the provider supporting its decision to not grant an exemption.

(2) The health care provider may make such a request at
any time, but it may make not more than one such request for the
same service, device, or drug in a calendar year.

(3) The department or its designee shall comply with such
a request.

(C) A health care provider may appeal the department or
its designee's decision to deny an exemption.

(D) The department or its designee shall not require a
health care provider to request an exemption provided under
division (A) of this section.

(E) When an exemption is granted under division (A) of
this section for a health care service, medical device, or drug,
the department or its designee shall notify the health care
provider in question. The notice shall include all of the following information:

(1) A statement that the health care provider qualifies for an exemption to a prior authorization requirement;

(2) The health care service, medical device, or drug to which the exemption applies;

(3) The dates the exemption will begin and end.

(F)(1) At the end of the twelve-month exemption period, the department or its designee may evaluate an exemption it has granted under division (A) of this section.

(2)(a) When conducting such an evaluation, the department or its designee shall review ten claims submitted to the department or its designee, selected at random, for the health care service, medical device, or drug in question.

(b) The reviewed claims shall be from the immediately preceding three months. If there are not ten relevant claims in the preceding three months, the department or its designee may review earlier claims.

(3)(a) If less than eighty per cent of the claims reviewed would have been approved based on medical necessity, then the department or its designee may revoke the exemption provided under division (A) of this section.

(b) If the department or its designee revokes an exemption, it shall provide the health care provider with both of the following:

(i) The information it relied upon in making its determination;
(ii) A plain language explanation of how to appeal the decision.

(4) The department or its designee shall not evaluate a health care provider's exemption relating to a particular service, device, or drug more than once every twelve months.

(5) Nothing in this section shall be construed as requiring the department or its designee to evaluate an existing exemption.

(G) If an exemption is revoked and not appealed, the exemption shall remain in effect until thirty days after the date the department or its designee notifies the health care provider of the department or its designee's decision to revoke the exemption.

(H) A health care provider may appeal the revocation of an exemption within thirty days of receiving notice of the revocation. If the health care provider appeals the revocation and the revocation is upheld, the exemption remains in effect until five days after the date the revocation is upheld.

(I) A decision to revoke or deny an exemption shall only be made by a health care provider licensed in this state who practices the same or a similar specialty as the health care provider being considered for an exemption and who has experience in providing the service, device, or drug to which the exemption or potential exemption applies.

(J) Nothing in this section shall be construed as prohibiting the department or its designee from making an administrative denial of a claim.

Section 2. That existing section 5160.34 of the Revised Code is hereby repealed.