9149

IN ASSEMBLY

January 31, 2022

Introduced by M. of A. HUNTER -- read once and referred to the Committee on Insurance

AN ACT to amend the insurance law and the social services law, in relation to requiring health insurance policies and medicaid to cover biomarker testing for certain purposes

The People of the State of New York, represented in Senate and Assembly, do enact as follows:

Section 1. Subsection (i) of section 3216 of the insurance law is 1 amended by adding a new paragraph 11-b to read as follows: 2 (11-b) (A) Every policy which provides medical, major medical, or 3 4 similar comprehensive-type coverage shall provide coverage for biomarker 5 testing for the purposes of diagnosis, treatment, appropriate manage-6 ment, or ongoing monitoring of a covered person's disease or condition 7 when the test is supported by medical and scientific evidence, includ-8 ing, but not limited to: 9 (i) labeled indications for a test approved or cleared by the food and 10 drug administration of the United States government or indicated tests for a food and drug administration approved drug; 11 12 (ii) centers for medicare and medicaid services national coverage determinations and medicare administrative contractor local coverage 13 14 determinations; or 15 (iii) nationally recognized clinical practice guidelines and consensus 16 statements. (B) Such coverage shall be provided in a manner that shall limit 17 18 disruptions in care including the need for multiple biopsies or biospe-19 cimen samples. 20 (C) The covered person and prescribing practitioner shall have access 21 to a clear, readily accessible, and convenient process to request an 22 exception to a coverage policy provided pursuant to the provisions of 23 this paragraph. Such process shall be made readily accessible on the 24 website of the insurer. (D) As used in this paragraph, the following terms shall have the 25

26 following meanings:

EXPLANATION--Matter in <u>italics</u> (underscored) is new; matter in brackets [-] is old law to be omitted.

LBD13311-02-2

A. 9149

1	(i) "Biomarker" means a characteristic that is objectively measured
2	and evaluated as an indicator of normal biological processes, pathogenic
3	processes, or pharmacologic responses to a specific therapeutic inter-
4	vention. Biomarkers include but are not limited to gene mutations or
5	protein expression.
6	(ii) "Biomarker testing" means the analysis of a patient's tissue,
7	blood, or other biospecimen for the presence of a biomarker. Biomarker
8	testing includes but is not limited to single-analyte tests, multi-plex
9	panel tests, and whole genome sequencing.
10	(iii) "Consensus statements" means statements developed by an inde-
11	pendent, multidisciplinary panel of experts utilizing a transparent
12^{11}	methodology and reporting structure and with a conflict of interest
13	policy. Such statements are aimed at specific clinical circumstances and
14^{13}	base the statements on the best available evidence for the purpose of
$14 \\ 15$	optimizing the outcomes of clinical care.
16	(iv) "Nationally recognized clinical practice guidelines" means
17	evidence-based clinical practice guidelines developed by independent
18	organizations or medical professional societies utilizing a transparent
19	methodology and reporting structure and with a conflict of interest
20	policy. Clinical practice guidelines establish standards of care
21	informed by a systematic review of evidence and an assessment of the
22	benefits and costs of alternative care options and include recommenda-
23	tions intended to optimize patient care.
24	§ 2. Subsection (1) of section 3221 of the insurance law is amended by
25	adding a new paragraph 11-b to read as follows:
26	(11-b) (A) Every insurer delivering a group or blanket policy or issu-
27	ing a group or blanket policy for delivery in this state that provides
28	coverage for medical, major medical, or similar comprehensive-type
29	coverage shall provide coverage for biomarker testing for the purposes
30	of diagnosis, treatment, appropriate management, or ongoing monitoring
31	of a covered person's disease or condition when the test is supported by
32	medical and scientific evidence, including, but not limited to:
33	(i) labeled indications for a test approved or cleared by the food and
34	drug administration of the United States government or indicated tests
35	for a food and drug administration approved drug;
36	(ii) centers for medicare and medicaid services national coverage
37	determinations and medicare administrative contractor local coverage
38	determinations; or
39	(iii) nationally recognized clinical practice guidelines and consensus
40	statements.
41	(B) Such coverage shall be provided in a manner that shall limit
42	disruptions in care including the need for multiple biopsies or biospe-
43	<u>cimen samples.</u>
44	(C) The covered person and prescribing practitioner shall have access
45	to a clear, readily accessible, and convenient process to request an
46	exception to a coverage policy provided pursuant to the provisions of
47	this paragraph. Such process shall be made readily accessible on the
48	website of the insurer.
49	(D) As used in this paragraph, the following terms shall have the
50	following meanings:
51	(i) "Biomarker" means a characteristic that is objectively measured
52	and evaluated as an indicator of normal biological processes, pathogenic
53	processes, or pharmacologic responses to a specific therapeutic inter-
54	vention. Biomarkers include but are not limited to gene mutations or

55 protein expression.

A. 9149

1	(ii) "Dispersion togeting" means the applying of a patientic tiggue
1	(ii) "Biomarker testing" means the analysis of a patient's tissue,
2	blood, or other biospecimen for the presence of a biomarker. Biomarker
3	testing includes but is not limited to single-analyte tests, multi-plex
4 5	panel tests, and whole genome sequencing.
	(iii) "Consensus statements" means statements developed by an inde-
6	pendent, multidisciplinary panel of experts utilizing a transparent
7	methodology and reporting structure and with a conflict of interest
8	policy. Such statements are aimed at specific clinical circumstances and
9	base the statements on the best available evidence for the purpose of
10	optimizing the outcomes of clinical care.
11	(iv) "Nationally recognized clinical practice guidelines" means
12	evidence-based clinical practice guidelines developed by independent
13 14	organizations or medical professional societies utilizing a transparent methodology and reporting structure and with a conflict of interest
14 15	policy. Clinical practice quidelines establish standards of care
15 16	informed by a systematic review of evidence and an assessment of the
17	benefits and costs of alternative care options and include recommenda-
18 19	<pre>tions intended to optimize patient care. § 3. Section 4303 of the insurance law is amended by adding a new</pre>
20	
20 21	<pre>subsection (p-1) to read as follows: (p-1) (1) A medical expense indemnity corporation, a hospital service</pre>
22	corporation or a health service corporation that provides coverage for
23	medical, major medical, or similar comprehensive-type coverage shall
23 24	provide coverage for biomarker testing for the purposes of diagnosis,
25	treatment, appropriate management, or ongoing monitoring of a covered
26	person's disease or condition when the test is supported by medical and
27	scientific evidence, including, but not limited to:
28	(A) labeled indications for a test approved or cleared by the food and
29	drug administration of the United States government or indicated tests
30	for a food and drug administration approved drug;
31	(B) centers for medicare and medicaid services national coverage
32	determinations and medicare administrative contractor local coverage
33	determinations; or
34	(C) nationally recognized clinical practice guidelines and consensus
35	statements.
36	(2) Such coverage shall be provided in a manner that shall limit
37	disruptions in care including the need for multiple biopsies or biospe-
38	cimen samples.
39	(3) The covered person and prescribing practitioner shall have access
40	to a clear, readily accessible, and convenient process to request an
41	exception to a coverage policy provided pursuant to the provisions of
42	this subsection. Such process shall be made readily accessible on the
43	website of the insurer.
44	(4) As used in this subsection, the following terms shall have the
45	following meanings:
46	(A) "Biomarker" means a characteristic that is objectively measured
47	and evaluated as an indicator of normal biological processes, pathogenic
48	processes, or pharmacologic responses to a specific therapeutic inter-
49	vention. Biomarkers include but are not limited to gene mutations or
50	
00	protein expression.
51	protein expression. (B) "Biomarker testing" means the analysis of a patient's tissue,
	(B) "Biomarker testing" means the analysis of a patient's tissue, blood, or other biospecimen for the presence of a biomarker. Biomarker
51	(B) "Biomarker testing" means the analysis of a patient's tissue,
51 52	(B) "Biomarker testing" means the analysis of a patient's tissue, blood, or other biospecimen for the presence of a biomarker. Biomarker

56 ent, multidisciplinary panel of experts utilizing a transparent method-

A. 9149

1	ology and reporting structure and with a conflict of interest policy.
2	Such statements are aimed at specific clinical circumstances and base
3	the statements on the best available evidence for the purpose of opti-
4	mizing the outcomes of clinical care.
5	(D) "Nationally recognized clinical practice guidelines" means
б	evidence-based clinical practice guidelines developed by independent
7	organizations or medical professional societies utilizing a transparent
8	methodology and reporting structure and with a conflict of interest
9	policy. Clinical practice guidelines establish standards of care
10	informed by a systematic review of evidence and an assessment of the
11	benefits and costs of alternative care options and include recommenda-
12	tions intended to optimize patient care.
13	§ 4. Subdivision 2 of section 365-a of the social services law is
14	amended by adding a new paragraph (jj) to read as follows:
15	(jj) (i) biomarker testing for the purposes of diagnosis, treatment,
16	appropriate management, or ongoing monitoring of a recipient's disease
17	or condition when the test is supported by medical and scientific
18	evidence, including, but not limited to:
19	(1) labeled indications for a test approved or cleared by the food and
20	drug administration of the United States government or indicated tests
21	for a food and drug administration approved drug;
22	(2) centers for medicare and medicaid services national coverage
23	determinations and medicare administrative contractor local coverage
24	determinations; or
25	(3) nationally recognized clinical practice guidelines and consensus
26	statements.
27	(ii) Risk-bearing entities contracted to the medicaid program to
28	deliver services to recipients shall provide biomarker testing at the
29	same scope, duration and frequency as the medicaid program otherwise
30	provides to enrollees.
31	(iii) The recipient and participating provider shall have access to a
32	clear, readily accessible, and convenient process to request an excep-
33	tion to a coverage policy of the medicaid program or by risk-bearing
34	entities contracted to the medicaid program. Such process shall be made
35	readily accessible to all participating providers and enrollees online.
36	(iv) As used in this paragraph, the following terms shall have the
37	following meanings:
38	(1) "Biomarker" means a characteristic that is objectively measured
39	and evaluated as an indicator of normal biological processes, pathogenic
40	processes, or pharmacologic responses to a specific therapeutic inter-
41	vention. Biomarkers include but are not limited to gene mutations or
42	protein expression.
43	(2) "Biomarker testing" means the analysis of a patient's tissue,
44	blood, or other biospecimen for the presence of a biomarker. Biomarker
45	testing includes but is not limited to single-analyte tests, multi-plex
46	panel tests, and whole genome sequencing.
47	(3) "Consensus statements" means statements developed by an independ-
48	ent, multidisciplinary panel of experts utilizing a transparent method-
49	ology and reporting structure and with a conflict of interest policy.
50	Such statements are aimed at specific clinical circumstances and base
51	the statements on the best available evidence for the purpose of opti-
52	mizing the outcomes of clinical care.
53 E4	(4) "Nationally recognized clinical practice guidelines" means
54 55	evidence-based clinical practice guidelines developed by independent
55	
56	organizations or medical professional societies utilizing a transparent methodology and reporting structure and with a conflict of interest

1	policy. Clinical practice guidelines establish standards of care
2	informed by a systematic review of evidence and an assessment of the
3	benefits and costs of alternative care options and include recommenda-
4	tions intended to optimize patient care.
5	§ 5. This act shall take effect January 1, 2023 and shall apply to all
6	policies and contracts issued, renewed, modified, altered or amended on
7	or after such date.