### **101ST GENERAL ASSEMBLY**

# State of Illinois

# 2019 and 2020

#### HB4314

Introduced 1/29/2020, by Rep. Camille Y. Lilly

## SYNOPSIS AS INTRODUCED:

305 ILCS 5/5-5.27 new

Amends the Medical Assistance Article of the Illinois Public Aid Code. Provides that the medical assistance program shall provide coverage for routine care costs that are incurred in the course of an approved clinical trial if the medical assistance program would provide coverage for the same routine care costs not incurred in a clinical trial. Defines "approved clinical trial" to mean a phase I, II, III, or IV clinical trial involving the prevention, detection, or treatment of cancer or any other life-threatening disease or condition. Defines "routine care cost" to mean the cost of medically necessary services related to the care method that is under evaluation in a clinical trial, including the cost of services related to the detection and treatment of any complications arising from the patient's medical care and any complications related to participation in the clinical trial. Defines other terms.

LRB101 15175 KTG 65493 b

FISCAL NOTE ACT MAY APPLY

A BILL FOR

HB4314

1

AN ACT concerning public aid.

# 2 Be it enacted by the People of the State of Illinois, 3 represented in the General Assembly:

4 Section 5. The Illinois Public Aid Code is amended by 5 changing Section 5-5.27 as follows:

(305 ILCS 5/5-5.27 new) 6 7 Sec. 5-5.27. Coverage for clinical trials. (a) The medical assistance program shall provide coverage 8 9 for routine care costs that are incurred in the course of an 10 approved clinical trial if the medical assistance program would provide coverage for the same routine care costs not incurred 11 12 in a clinical trial. (b) The coverage that must be provided under this Section 13 14 is subject to the terms, conditions, restrictions, exclusions, and limitations that apply generally under the medical 15 assistance program, including terms, conditions, restrictions, 16 exclusions, or limitations that apply to health care services 17 rendered by participating providers and nonparticipating 18 19 providers. 20 (c) As used in this Section: 21 "Approved clinical trial" means a phase I, II, III, or IV 22 clinical trial involving the prevention, detection, or

23 treatment of cancer or any other life-threatening disease or

1	condition if one or more of the following conditions apply:					
2	(1) the Department makes a determination that the study					
3	or investigation is an approved clinical trial;					
4	(2) the study or investigation is conducted under an					
5	investigational new drug application or an investigational					
6	device exemption reviewed by the federal Food and Drug					
7	Administration;					
8	(3) the study or investigation is a drug trial that is					
9	exempt from having an investigational new drug application					
10	or an investigational device exemption from the federal					
11	Food and Drug Administration; or					
12	(4) the study or investigation is approved or funded					
13	(which may include funding through in-kind contributions)					
14	by:					
15	(A) the National Institutes of Health;					
16	(B) the Centers for Disease Control and					
17	Prevention;					
18	(C) the Agency for Healthcare Research and					
19	Quality;					
20	(D) the Patient-Centered Outcomes Research					
21	Institute;					
22	(E) the federal Centers for Medicare and Medicaid					
23	Services;					
24	(F) a cooperative group or center of any of the					
25	entities described in subparagraphs (A) through (E) or					
26	the United States Department of Defense or the United					

1	States Department of Veterans Affairs;					
2	(G) a qualified non-governmental research entity					
3	identified in the guidelines issued by the National					
4	Institutes of Health for center support grants; or					
5	(H) the United States Department of Veterans					
6	Affairs, the United States Department of Defense, or					
7	the United States Department of Energy, provided that					
8	review and approval of the study or investigation					
9	occurs through a system of peer review that is					
10	comparable to the peer review of studies performed by					
11	the National Institutes of Health, including an					
12	unbiased review of the highest scientific standards by					
13	qualified individuals who have no interest in the					
14	outcome of the review.					
15	"Care method" means the use of a particular drug or device					
16	in a particular manner.					
17	"Life-threatening disease or condition" means a disease or					
18	condition from which the likelihood of death is probable unless					
19	the course of the disease or condition is interrupted.					
20	"Routine care cost" means the cost of medically necessary					
21	services related to the care method that is under evaluation in					
22	a clinical trial. Routine care costs include the cost of					
23	services related to the detection and treatment of any					
24	complications arising from the patient's medical care,					
25	including any complications related to participation in the					
26	clinical trial. The term does not include the following:					

	HB4314		- 4 -	LRB101	15175 KTG 65493 b				
1	(1)	The drug or	device that	<u>is under</u>	evaluation in a				
2	<u>clinica</u>	clinical trial.							
3	(2)	Items or ser	vices that are	<u>.</u>					
4		(A) provide	ed solely fo	or data	collection and				
5	ana	lysis and not	t in the dired	ct clinic	al management of				
6	an	<u>individual en</u>	rolled in a cl	inical ti	rial;				
7		<u>(B) customar</u>	cily provided	at no co	st by a research				
8	spc	nsor to an in	dividual enro	lled in a	a clinical trial;				
9	or								
10		(C) provided	l solely to de	termine e	eligibility of an				
11	ind	lividual for p	articipation :	in a clin	ical trial.				