

THE GENERAL ASSEMBLY OF PENNSYLVANIA

HOUSE BILL

No. 508 Session of 2011

INTRODUCED BY DeLUCA, DEASY, FABRIZIO, D. COSTA, CALTAGIRONE, BOBACK, BUXTON, FLECK, HARKINS, JOSEPHS, GEORGE, KOTIK, MOUL, MURT, GOODMAN, GRELL, HARRIS, HESS, HORNAMAN, KAVULICH, KIRKLAND, M. O'BRIEN, PASHINSKI, SCAVELLO, K. SMITH, SONNEY, STABACK, STURLA, WHITE AND YOUNGBLOOD, FEBRUARY 4, 2011

REFERRED TO COMMITTEE ON INSURANCE, FEBRUARY 4, 2011

AN ACT

1 Providing for insurance coverage for patient costs associated
2 with cancer clinical trials.

3 The General Assembly of the Commonwealth of Pennsylvania
4 hereby enacts as follows:

5 Section 1. Short title.

6 This act shall be known and may be cited as the Cancer
7 Clinical Trials Act.

8 Section 2. Definitions.

9 The following words and phrases when used in this act shall
10 have the meanings given to them in this section unless the
11 context clearly indicates otherwise:

12 "Carrier." An insurance company, health service corporation,
13 hospital service corporation, medical service corporation or
14 health maintenance organization authorized to issue health
15 benefits plans in this Commonwealth.

16 "Cooperative group." A formal network of facilities that

1 collaborates on research projects and that has an established
2 National Institutes of Health approved peer review program
3 operating within the group, including the National Cancer
4 Institute clinical cooperative group and the National Cancer
5 Institute community clinical oncology program.

6 "Health benefits plan." A hospital and medical expense
7 insurance policy or certificate; health, hospital or medical
8 service corporation contract or certificate; or health
9 maintenance organization subscriber contract or certificate
10 delivered or issued for delivery in this Commonwealth by any
11 carrier. The term excludes the following plans, policies or
12 contracts: specified disease, CHAMPUS supplement, accident only,
13 credit, disability, long-term care, coverage for Medicare
14 services pursuant to a contract with the Federal Government,
15 Medicare supplement, dental only or vision only, insurance
16 issued as a supplement to liability insurance, coverage arising
17 out of a workers' compensation or similar law, hospital
18 confinement or other supplemental limited benefit insurance
19 coverage or automobile medical payment insurance.

20 "Institutional review board." Any board, committee or other
21 group that is both:

22 (1) Formally designated by an institution to approve the
23 initiation of and to conduct periodic review of biomedical
24 research involving human subjects and in which the primary
25 purpose of such review is to assure the protection of the
26 rights and welfare of the human subjects and not to review a
27 clinical trial for scientific merit.

28 (2) Approved by the National Institutes of Health office
29 for protection from research risks.

30 "Multiple project assurance contract." A contract between an

1 institution and the United States Department of Health and Human
2 Services that defines the relationship of the institution to the
3 United States Department of Health and Human Services and that
4 sets out the responsibilities of the institution and the
5 procedures that will be used by the institution to protect human
6 subjects.

7 "Patient." The subscriber, insured or enrollee or the
8 covered dependent of the subscriber, insured or enrollee.

9 "Routine care costs." Physician fees, laboratory expenses
10 and expenses associated with the hospitalization, administering
11 of treatment and evaluation of the patient during the course of
12 treatment which are consistent with usual and customary patterns
13 and standards of care incurred whenever an enrollee, subscriber
14 or insured receives medical care associated with an approved
15 cancer clinical trial and which would be covered if such items
16 and services were provided other than in connection with an
17 approved cancer clinical trial.

18 Section 3. Coverage for clinical cancer trials.

19 (a) General rule.--A carrier is not obligated to pay any
20 costs, other than routine care costs, that are directly
21 associated with a cancer clinical trial that is offered in this
22 Commonwealth and in which the subscriber, insured or enrollee
23 participates voluntarily. A cancer clinical trial is a course of
24 treatment in which all of the following apply:

25 (1) The treatment is part of a scientific study of a new
26 therapy or intervention that is being conducted at an
27 institution in this Commonwealth, that is for the treatment,
28 palliation or prevention of cancer in humans and in which the
29 scientific study includes all of the following:

30 (i) Specific goals.

1 (ii) A rationale and background for the study.

2 (iii) Criteria for patient selection.

3 (iv) Specific directions for administering the
4 therapy and monitoring patients.

5 (v) A definition of quantitative measures for
6 determining treatment response.

7 (vi) Methods for documenting and treating adverse
8 reactions.

9 (2) The treatment is being provided as part of a study
10 being conducted in a Phase I, Phase II, Phase III or Phase IV
11 cancer clinical trial.

12 (3) The treatment is being provided as part of a study
13 being conducted in accordance with a clinical trial approved
14 by at least one of the following:

15 (i) One of the National Institutes of Health.

16 (ii) A National Institutes of Health cooperative
17 group or center.

18 (iii) The United States Food and Drug Administration
19 in the form of an investigational new drug application.

20 (iv) The United States Department of Defense.

21 (v) The United States Department of Veterans
22 Affairs.

23 (vi) A qualified research entity that meets the
24 criteria established by the National Institutes of Health
25 for grant eligibility.

26 (vii) A panel of qualified recognized experts in
27 clinical research within academic health institutions in
28 this Commonwealth.

29 (4) The proposed treatment or study has been reviewed
30 and approved by an institutional review board of an

1 institution in this Commonwealth.

2 (5) The personnel providing the treatment or conducting
3 the study:

4 (i) Are providing the treatment or conducting the
5 study within their scope of practice, experience and
6 training and are capable of providing the treatment
7 because of their experience, training and volume of
8 patients treated to maintain expertise.

9 (ii) Agree to accept reimbursement as payment in
10 full from the carrier at the rates that are established
11 by the carrier and that are not more than the level of
12 reimbursement applicable to other similar services
13 provided by health care providers with the carrier's
14 provider network.

15 (6) There is no clearly superior, noninvestigational
16 treatment alternative.

17 (7) The available clinical or preclinical data provide a
18 reasonable expectation that the treatment will be at least as
19 efficacious as any noninvestigational alternative.

20 (b) Liability.--Pursuant to the patient informed consent
21 document, no party is liable for damages associated with the
22 treatment provided during any phase of a cancer clinical trial.

23 (c) Benefits.--Each health benefits plan delivered or issued
24 for delivery in this Commonwealth shall provide benefits under
25 the plan, and those benefits shall not supplant any portion of
26 the clinical trial that is customarily paid for by government,
27 biotechnical, pharmaceutical or medical device industry sources.

28 (d) Remedy.--This section does not create any private right
29 or cause of action for or on behalf of any patient against the
30 carrier. This section provides solely an administrative remedy

1 for any violation of this section or any related rule.

2 (e) Deductibles and other cost sharing.--Nothing in this
3 section prohibits the carrier from imposing deductibles,
4 coinsurance or other cost sharing measures in relation to
5 benefits provided pursuant to this section.

6 Section 4. Applicability.

7 This act applies to health benefit plans issued or renewed on
8 or after January 1, 2012.

9 Section 5. Effective date.

10 This act shall take effect immediately.