

1 State of Arkansas
2 89th General Assembly
3 Regular Session, 2013
4

As Engrossed: H2/28/13

A Bill

HOUSE BILL 1267

5 By: Representatives Kerr, Branscum, Carnine, Clemmer, Cozart, Ferguson, Gillam, Gossage, Hobbs,
6 Leding, Linck, Lowery, Magie, Sabin
7 By: Senators J. Dismang, Holland, J. Hutchinson, J. Key, Rapert
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For An Act To Be Entitled

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10 AN ACT TO REQUIRE A PRIOR APPROVAL PROCESS FOR
11 EXPERIMENTAL AND INVESTIGATIONAL SURGICAL PROCEDURES
12 AND MEDICAL DEVICES; AND FOR OTHER PURPOSES.
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Subtitle

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16 AN ACT TO REQUIRE A PRIOR APPROVAL
17 PROCESS FOR EXPERIMENTAL AND
18 INVESTIGATIONAL SURGICAL PROCEDURES AND
19 MEDICAL DEVICES.
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22 BE IT ENACTED BY THE GENERAL ASSEMBLY OF THE STATE OF ARKANSAS:
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24 SECTION 1. Arkansas Code Title 23, Chapter 86, Subchapter 1, is
25 amended to add an additional section to read as follows:

26 23-86-122. Prior approval process for experimental and investigational
27 surgical products and medical devices.

28 (a) "Health carrier" means a health maintenance organization, hospital
29 medical service corporation, or a disability insurance company.

30 (A) "Health carrier" includes a self-insured governmental
31 or church plan and third-party administrators that administer or adjust
32 disability benefits for a disability insurer, hospital medical service
33 corporation, health maintenance organization, self-insured governmental plan
34 or self-insured church plan.

35 (B) "Health carrier" does not include:

36 (i) An automobile insurer paying medical or hospital



1 benefits under § 23-89-202(1) or a self-insured employer health benefits
2 plan; or

3 (ii) A person, company, or organization licensed or
4 registered to issue or who issues any insurance policy or insurance contract
5 in this state as described in §§ 23-62-102 and 23-62-104 – 23-62-107
6 providing medical or hospital benefits for accidental injury or disability.

7 (b) A health carrier that denies coverage for a specific surgical
8 product or medical device approved for marketing by the United States Food
9 and Drug Administration as experimental, investigational, or both shall
10 develop a process by which a surgeon, before utilizing the device or
11 treatment, may present medical evidence to obtain a review for the individual
12 patient for coverage of the surgical product or medical device.

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14 /s/Kerr
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