2012 SESSION

[S 544]

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VIRGINIA ACTS OF ASSEMBLY - CHAPTER

An Act to amend and reenact § 32.1-229 of the Code of Virginia, relating to mammogram reports. 2

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Approved

5 Be it enacted by the General Assembly of Virginia:

- 1. That § 32.1-229 of the Code of Virginia is amended and reenacted as follows: 6 7
 - § 32.1-229. Powers and duties of the Board.
 - The Board shall:

9 1. Establish a program of effective regulation of sources of radiation for the protection of the public 10 health and safety, including a program of education and technical assistance relating to radon that is targeted to those areas of the Commonwealth known to have high radon levels. 11

12 2. Establish a program to promote the orderly regulation of radiation within the Commonwealth, among the states and between the federal government and the Commonwealth and to facilitate 13 intergovernmental cooperation with respect to use and regulation of sources of radiation to the end that 14 15 duplication of regulation may be minimized.

3. Establish a program to permit maximum utilization of sources of radiation consistent with the 16 17 public health and safety.

4. Promulgate regulations providing for (i) general or specific licenses to use, manufacture, produce, 18 19 transfer, receive, acquire, own or possess quantities of, or devices or equipment utilizing, by-product, 20 source, special nuclear materials, or other radioactive material occurring naturally or produced 21 artificially, (ii) registration of the possession of a source of radiation and of information with respect 22 thereto, and (iii) regulation of by-product, source and special nuclear material.

23 5. Encourage, participate in and conduct studies, investigations, training, research and demonstrations 24 relating to control of sources of radiation.

6. Establish fee schedules for the licensure of radioactive materials.

26 7. Establish guidelines to require the licensed facilities or physicians' offices where mammography 27 services are performed to offer to the patient, prior to departure, development of such films to ensure integrity and quality of the film. When film developing is not available or the patient chooses not to 28 29 wait, the patient shall be notified within two business days if another mammogram is necessary. This 30 requirement does not imply or require that a diagnostic opinion be made at the time of the mammogram. 31 The interpreting physician may require that the mammogram be retaken if, in the opinion of the 32 physician, the study is of inadequate quality. Such guidelines shall also require the licensed facility or 33 physician's office where mammography services are performed to (i) include information on breast 34 density in mammogram letters sent to patients pursuant to regulations implementing the Mammography Quality Standards Act promulgated by the U.S. Food and Drug Administration, and (ii) include in 35 letters sent to patients who have dense breast tissue, as determined by the interpreting physician based 36 37 on standards promulgated by the American College of Radiology, the following notice:

38 "YOUR MAMMÖGRAM DEMONSTRATES THÅT YOU MAY HAVE DENSE BREAST TISSUE, 39 WHICH CAN HIDE CANCER OR OTHER ABNORMALITIES. A REPORT OF YOUR 40 MAMMOGRAPHY RESULTS, WHICH CONTAINS INFORMATION ABOUT YOUR BREAST DENSITY, 41 HAS BEEN SENT TO YOUR REFERRING PHYSICIAN'S OFFICE, AND YOU SHOULD CONTACT 42 YOUR PHYSICIAN IF YOU HAVE ANY OUESTIONS OR CONCERNS ABOUT THIS REPORT."

43 8. Issue such orders or modifications thereof as may be necessary in connection with proceedings 44 under this title.

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