

1 A bill to be entitled
 2 An act relating to clinical laboratories; amending s.
 3 483.181, F.S.; requiring clinical laboratories to accept
 4 human specimens submitted by advanced registered nurse
 5 practitioners; providing an effective date.

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 7 Be It Enacted by the Legislature of the State of Florida:

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 9 Section 1. Section 483.181, Florida Statutes, is amended
 10 to read:

11 483.181 Acceptance, collection, identification, and
 12 examination of specimens.--

13 (1) A clinical laboratory may examine human specimens at
 14 the request only of a licensed practitioner or other person
 15 authorized by law to use the findings of clinical laboratory
 16 examinations. An individual forwarding a sample of the
 17 individual's own blood to a clinical laboratory, when such blood
 18 sample has been taken pursuant to a home access HIV test kit
 19 approved by the United States Food and Drug Administration,
 20 shall be considered a person authorized to request and use a
 21 clinical laboratory test for human immunodeficiency virus, for
 22 the purposes of this part.

23 (2) The results of a test must be reported directly to the
 24 licensed practitioner or other authorized person who requested
 25 it. The report must include the name and address of the clinical
 26 laboratory in which the test was actually performed, unless the
 27 test was performed in a hospital laboratory and the report
 28 becomes an integral part of the hospital record.

29 (3) The results of clinical laboratory tests performed by
30 a clinical laboratory complying with this part and performed
31 before a patient's admission to a facility licensed under
32 chapter 395 must be accepted in lieu of clinical laboratory
33 tests required upon a patient's admission to the facility and in
34 lieu of tests that may be ordered for patients of the facility,
35 except that the facility may not be required to accept
36 transfusion compatibility test results. The agency shall
37 establish, by rule, standards for accepting laboratory test
38 results to specify acceptable timeframes for such laboratory
39 tests to assure that the timeframes do not adversely affect the
40 accuracy of the test.

41 (4) All specimens accepted by a clinical laboratory must
42 be tested on the premises, except that specimens for
43 infrequently performed tests may be forwarded for examination to
44 another clinical laboratory approved under this part. This
45 subsection does not prohibit referring specimens to a clinical
46 laboratory excepted under s. 483.031. However, the clinical
47 laboratory director of the referring clinical laboratory must
48 assume complete responsibility.

49 (5) A clinical laboratory licensed under this part must
50 accept a human specimen submitted for examination by a
51 practitioner licensed under chapter 458, chapter 459, chapter
52 460, chapter 461, chapter 462, s. 464.012, or chapter 466, if
53 the specimen and test are the type performed by the clinical
54 laboratory. A clinical laboratory may only refuse a specimen
55 based upon a history of nonpayment for services by the
56 practitioner. A clinical laboratory shall not charge different

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57 | prices for tests based upon the chapter under which a
58 | practitioner submitting a specimen for testing is licensed.

59 | Section 2. This act shall take effect July 1, 2008.