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House Bill 966 (AS PASSED HOUSE AND SENATE)

By: Representatives Cooper of the 43rd, Oliver of the 82nd, Rutledge of the 109th, Watson of the 166th, Weldon of the 3rd, and others

A BILL TO BE ENTITLED AN ACT

1 To amend Chapter 2A of Title 31 of the Official Code of Georgia Annotated, relating to the

- 2 Department of Public Health, so as to establish within the Department of Public Health the
- 3 Alzheimer's Disease Registry; to provide for the purpose of the registry; to provide for
- 4 promulgation and criteria of rules; to provide for confidentiality of data; to provide for
- 5 compliance with P. L. 104-191, the federal Health Insurance Portability and Accountability
- 6 Act of 1996; to provide for related matters; to repeal conflicting laws; and for other purposes.

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BE IT ENACTED BY THE GENERAL ASSEMBLY OF GEORGIA:

9 **SECTION 1.**

- 10 Chapter 2A of Title 31 of the Official Code of Georgia Annotated, relating to the Department
- of Public Health, is amended by adding a new Code section to read as follows:
- 12 <u>"31-2A-16.</u>
- 13 (a) There is established within the Department of Public Health the Alzheimer's Disease
- 14 Registry.
- 15 (b) The purpose of the registry shall be to assist in the development of public policy and
- planning relative to Alzheimer's disease and related disorders. The registry shall provide
- 17 <u>a central data base of individuals with Alzheimer's disease or related disorders.</u>
- 18 (c) The department shall establish procedures and promulgate rules and regulations for the
- 19 <u>establishment and operation of the registry. Such procedures, rules, and regulations shall</u>
- 20 provide for:
- 21 (1) Collecting and evaluating data regarding the prevalence of Alzheimer's disease and
- 22 related disorders in Georgia, including who shall report the data to the registry;
- 23 (2) Determining what information shall be maintained in the registry and the length of
- 24 time such data shall be available;
- 25 (3) Sharing of data for policy planning purposes;
- 26 (4) Disclosing nonidentifying data to support Alzheimer's and related disorder research;
- 27 (5) The methodology by which families and physicians of persons who are reported to
- 28 the registry shall be contacted to gather additional data; and

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29 <u>(</u>	(6)	Information about	public and	private resources.

- 30 (d) The collected data in the registry shall be confidential, and all persons to whom the
- 31 data is released shall maintain patient confidentiality. No publication of information,
- 32 <u>biotechnical research, or medical data shall be made that identifies any patient by name.</u>
- 33 The registry shall be established and regulated pursuant to the requirements of 42 U.S.C.
- 34 Section 1301, et seq., and P.L. 104-191, the federal Health Insurance Portability and
- 35 Accountability Act of 1996."

36 SECTION 2.

37 All laws and parts of laws in conflict with this Act are repealed.