## **HOUSE OF REPRESENTATIVES STAFF ANALYSIS**

BILL #: HB 17 CS Developmental Disabilities

**SPONSOR(S):** Kravitz & Others

TIED BILLS: IDEN./SIM. BILLS: CS/SB 428

REFERENCE	ACTION	ANALYST	STAFF DIRECTOR
1) Elder & Long-Term Care Committee	7 Y, 0 N, w/CS	Walsh	Liem
2) Health Care General Committee	9 Y, 0 N	Schiefelbein	Brown-Barrios
3) Health Care Appropriations Committee	11 Y, 0 N	Ekholm	Massengale
4) Health & Families Council	9 Y, 0 N	Walsh	Moore
5)			

## **SUMMARY ANALYSIS**

House Bill 17 CS amends section 409.912, Florida Statutes, related to the cost-effective purchasing of Medicaid services. The bill requires the Agency for Health Care Administration to develop and seek federal approval for a model home and community-based waiver to serve children who are diagnosed with familial dysautonomia.

The bill makes implementation of the waiver contingent upon the availability of funds and any limitations in the General Appropriations Act.

The bill takes effect July 1, 2005.

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### **FULL ANALYSIS**

## I. SUBSTANTIVE ANALYSIS

### A. HOUSE PRINCIPLES ANALYSIS:

**Provide limited government**—The bill creates Medicaid eligibility for a new group of people.

**Promote personal responsibility**—The bill creates an opportunity for families with children with familial dysautonomia to obtain Medicaid eligibility for their children by disregarding the family's income and considering the child as a "family of one" for determining income and assets.

**Empower families**—The bill creates Medicaid eligibility for a new group of people, increasing dependence of families upon government support or assistance.

### B. EFFECT OF PROPOSED CHANGES:

# **Background**

Familial dysautonomia (FD), or Riley-Day syndrome, is a genetic disease that is present from birth that results in incomplete development of the nervous system causing a decreased number of nerve cells. The affected nerve cells are those that control certain sensations and autonomic functions. The sensory nerve cells that are most severely affected are those responsible for pain, heat perception, and taste. The autonomic nerve cells control bodily functions such as sweating, swallowing, regulation of blood pressure and body temperature, and the ability to cry emotional tears. Secondary problems associated with the disease may include feeding problems, vomiting, poor growth, spinal curvature, and lung problems.

Familial dysautonomia patients can be expected to function independently if treatment is begun early and major disabilities are avoided. Affected individuals usually are of normal intelligence.

Familial dysautonomia is a recessive genetic disease, meaning that both parents carry the gene despite a lack of outward signs. The recessive gene causing FD occurs with a remarkably high carrier frequency in individuals of Eastern European Jewish ancestry (Ashkenazi Jewish extraction). It is estimated that one in 30 persons with Eastern European Jewish ancestry is a carrier of the FD gene. FD does not express itself in a consistent manner, and the costs for treatment vary depending upon the severity and types of symptoms experienced by the individual. Symptoms also vary by age.<sup>1</sup>

The Dysautonomia Foundation, Inc. in New York reports that, based on the FD world-wide registry, as of January 2004 there are over 340 people worldwide living with FD. One-third of them live in the metropolitan New York City area, one-third reside in Israel, and the remaining third live elsewhere in the United States and worldwide.<sup>2</sup> It has been reported that 17 persons with FD are residents of Florida.

The Department of Health, Children's Medical Services (CMS), currently provides services to children diagnosed with FD whose families meet certain income limitations. Under the federal requirements of OBRA 89 that Medicaid programs meet children's medical needs, CMS is able to provide care coordination for a range of needed services and therapies. However, CMS does not have funding to provide families of FD children with respite or behavioral services. Persons with FD are not currently served by the Agency for Persons with Disabilities (APD).<sup>3</sup>

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<sup>&</sup>lt;sup>1</sup> See, e.g., Jewish Genetic Diseases A Mazornet Guide at http://www.mazornet.com/genetics/familial\_dysautonomia.asp; Chicago Center for Jewish Genetic Disorders at http://www.jewishgeneticscenter.org/what/ashkenazi/familial2.asp.

<sup>&</sup>lt;sup>2</sup> Dysautonomia Foundation, Inc. at http://www.familialdysautonomia.org/history.htm

<sup>&</sup>lt;sup>3</sup> The Agency for Persons with Disabilities was formerly the Development Disabilities Program of the Department of Children and Families.

## **Medicaid Home and Community-based Waivers**

In 1981, Congress authorized the waiver of certain federal requirements to enable a state to provide home and community services (other than room and board) to individuals who would otherwise require institutional care reimbursed by Medicaid. The waiver programs are called "1915(c) waivers." Under 1915(c) waiver authority, states can provide services not usually covered by the Medicaid program, as long as these services are required to keep a person from being institutionalized.

The Model waiver (originally known as a Katie Beckett waiver) is a provision in the Tax Equity and Fiscal Responsibility Act (TEFRA 134) that was added to the Medicaid program in 1982. TEFRA 134 gives states the option to cover non-institutionalized children with disabilities. Before TEFRA 134 was enacted, if a child with disabilities lived at home, the parents' income and resources were automatically counted (deemed) as available for medical expenses. However, if the same child was institutionalized for 30 days or more, only the child's own income and resources were counted in determining financial eligibility.

TEFRA 134 amended the Medicaid law to give states the option to waive or disregard the deeming of parental income and resources for children under 18 years old who were living at home but who would otherwise be eligible for Medicaid-funded institutional care. Not counting parental income enables these children to receive Medicaid services at home or in other community settings. CMS reports that many states use this option. States must determine that:

- (1) The child requires the level of care provided in an institution.
- (2) It is appropriate to provide care outside the facility.
- (3) The cost of care at home is no more than the cost of institutional care. In states using this option, parents may choose either institutional or community care for their Medicaid eligible children.

## **Effect of Changes**

AHCA is directed to develop a model home and community-based waiver. If approved, Medicaid coverage would be available to children diagnosed with familial dysautonomia. It is likely that AHCA would serve children who are financially eligible by disregarding their parent's income and assets. Eligibility for the Medicaid waiver also provides access to all other Medicaid services.

## C. SECTION DIRECTORY:

Section 1: Amends s. 409.912, F. S., requiring AHCA to develop a model waiver program to serve children with familial dysautonomia.

**Section 2**: Provides an effective date.

# II. FISCAL ANALYSIS & ECONOMIC IMPACT STATEMENT

# A. FISCAL IMPACT ON STATE GOVERNMENT:

1. Revenues:

None.

## 2. Expenditures:

The fiscal impact of this bill is unknown as there is no historical cost data associated with serving persons with FD in either the Medicaid program or the Developmental Services program.

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There is an approved Medicaid Model Waiver that is for children under the age of 21 with certain medical conditions to help maintain them in their home instead of a hospital setting. The waiver services include case management and respite care. The average per child per month cost is \$1,330.

If the federal Centers for Medicare and Medicaid Services approves the model waiver authorized in the bill, children with FD who qualified for the waiver would be eligible not only for the waiver services but also for the entire package of Medicaid services.

## **B. FISCAL IMPACT ON LOCAL GOVERNMENTS:**

Revenues:

None.

2. Expenditures:

None.

#### C. DIRECT ECONOMIC IMPACT ON PRIVATE SECTOR:

Persons deemed eligible for the model waiver could receive the full array of Medicaid State Plan services, as well as the waiver services.

### D. FISCAL COMMENTS:

The bill provides that AHCA may implement the federally-approved waiver subject to the availability of funds and any limitations provided in the General Appropriations Act. Establishing Medicaid eligibility for the child under this waiver authority will also make the child eligible for all Medicaid services. However, Medicaid would coordinate benefits with any other third party payer as long as that payer was available. Very sick or disabled children frequently exhaust the lifetime cap on the family's health insurance benefits.

## III. COMMENTS

### A. CONSTITUTIONAL ISSUES:

1. Applicability of Municipality/County Mandates Provision:

This bill does not require counties or municipalities to spend funds or take an action requiring the expenditure of funds. This bill does not reduce the percentage of a state tax shared with counties or municipalities. This bill does not reduce the authority that municipalities have to raise revenue.

2. Other:

None.

### B. RULE-MAKING AUTHORITY:

AHCA is granted rulemaking authority to implement the provisions of the act.

C. DRAFTING ISSUES OR OTHER COMMENTS:

None.

# IV. AMENDMENTS/COMMITTEE SUBSTITUTE CHANGES

At its January 26, 2005, meeting, the Committee on Elder and Long Term Care adopted one amendment to the bill which deleted references to "Department of Children and Family Services." AHCA has

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responsibility for the existing model waiver program; neither DCF nor APD would administer the waiver if it receives federal approval.

This analysis is drafted to the Committee Substitute.

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