CHAPTER 266-A

INFORMATION AND REFERRAL SERVICES

§1641. Parkinson's Syndrome

The Department of Human Services shall establish, maintain and operate an information and referral service for Parkinson's Syndrome to assist in promoting the general health and welfare of Maine's citizens, including, but not limited to, the following specific purposes: [PL 1985, c. 737, Pt. A, §51 (NEW).]

1. Information. To provide educational materials to the medical community and other interested individuals relating to the nature and treatment of Parkinson's Syndrome; and [PL 1985, c. 737, Pt. A, §51 (NEW).]

2. Referral. To maintain a referral service to make available, upon request, the names, addresses and phone numbers, when known, of:
   A. Physicians who have an interest or expertise in Parkinson's Syndrome; and [PL 1985, c. 737, Pt. A, §51 (NEW).]
   B. Local or statewide support groups for Parkinson's Syndrome victims or their families and friends. [PL 1985, c. 737, Pt. A, §51 (NEW).]

SECTION HISTORY
PL 1985, c. 737, §A51 (NEW).

§1642. Down syndrome

The department shall establish, maintain and operate an information service for Down syndrome. For the purposes of this section, "Down syndrome" means a chromosomal condition caused by an error in cell division that results in the presence of an extra whole or partial copy of chromosome 21. [PL 2015, c. 269, §1 (NEW).]

1. Expectant or new parent; others. A hospital, physician, health care provider or certified nurse midwife who renders prenatal care or postnatal care or a genetic counselor who renders prenatal or postnatal genetic counseling shall, upon receipt of a positive test result from a prenatal or postnatal test for Down syndrome, offer the expectant or new parent information provided by the department under subsection 2. The department shall also make such information available to any other person who has received a positive test result from a prenatal or postnatal test for Down syndrome. [PL 2015, c. 269, §1 (NEW).]

2. Information provided. The department shall make available to a person who renders prenatal care, postnatal care or genetic counseling to expectant or new parents who receive a prenatal or postnatal diagnosis of Down syndrome the following:
   A. Up-to-date evidence-based written information about Down syndrome that includes physical, developmental, educational and psychosocial outcomes, life expectancy, clinical course and intellectual and functional development and treatment options. The information must have been reviewed by established medical experts in the field and national Down syndrome organizations; and [PL 2015, c. 269, §1 (NEW).]
   B. Contact information regarding support programs and services, including information hotlines specific to Down syndrome, resource centers and clearinghouses, national, statewide and local
Down syndrome organizations and other educational and support programs. [PL 2015, c. 269, §1 (NEW).]
[PL 2015, c. 269, §1 (NEW).]

3. Accessibility of information. Information provided under this section must be culturally and linguistically appropriate for a person receiving a positive prenatal diagnosis and for the family of a child receiving a postnatal diagnosis of Down syndrome.
[PL 2015, c. 269, §1 (NEW).]

SECTION HISTORY
PL 2015, c. 269, §1 (NEW).