

1                   A bill to be entitled  
2       An act relating to experimental treatments for  
3       terminal conditions and life-threatening rare  
4       diseases; providing a short title; amending s.  
5       499.0295, F.S.; providing legislative findings and  
6       intent; defining terms; authorizing licensed  
7       physicians to prescribe and eligible facilities to  
8       administer experimental treatments if certain  
9       requirements are met; requiring the department to  
10      adopt certain rules; requiring a specified written  
11      informed consent from eligible patients; requiring  
12      approval of a specified platform or master protocol by  
13      an institutional review board; requiring the  
14      Department of Health to establish and maintain a  
15      patient registry for specified purposes; requiring  
16      eligible facilities to submit certain information to  
17      the registry; requiring the department to enter into  
18      specified contracts with eligible facilities;  
19      providing a penalty for breach of such contact;  
20      requiring the department to annually publish a  
21      specified report; encouraging health insurers and  
22      health maintenance organizations to provide specified  
23      insurance coverage for experimental treatments;  
24      prohibiting such insurers and organizations from  
25      denying insurance coverage for experimental

26 treatments; providing construction; authorizing  
27 licensed physicians and eligible facilities to receive  
28 reimbursement for the prescription or administration  
29 of experimental treatments if certain requirements are  
30 met; establishing the reimbursement rate; requiring  
31 the department to adopt specified rules; providing for  
32 reimbursement rate expiration; requiring manufacturers  
33 of experimental treatments to provide for a specified  
34 temporary price concession set by the department;  
35 providing for expiration of such concession;  
36 authorizing the Department of Management Services to  
37 enter into a specified contract with an eligible  
38 facility or manufacturer; requiring the department to  
39 adopt a specified rule for such contract; requiring  
40 the Department of Health to adopt by rule certain  
41 procedures for licensing of experimental treatment  
42 centers; requiring the department to approve or deny a  
43 completed application within a specified timeframe;  
44 requiring the department to establish by rule certain  
45 procedural and operational standards; prohibiting  
46 certain actions by specified licensing boards and a  
47 specified state agency against a licensed physician in  
48 certain circumstances; providing construction and  
49 applicability; providing for limitation of liability;  
50 providing that certain participation is entirely

voluntary at all times; prohibiting public officials, public employees, and public agents from denying or attempting to deny access to experimental treatment; providing construction; requiring the department to adopt rules; providing an effective date.

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

Section 1. "This act may be cited as the "Promising Pathways Act."

**Section 2. Section 499.0295, Florida Statutes, is amended to read:**

(Substantial rewording of section. See s. 499.0295, F.S., for present text.)  
499.0295 Experimental treatments for terminal conditions and life-threatening rare diseases.—

(1) The Legislature finds that residents of this state with terminal conditions and life-threatening rare diseases may benefit from access to experimental treatments that have demonstrated initial human safety but are not yet approved by the United States Food and Drug Administration.

(2) It is the intent of the Legislature to:

(a) Create a safe, regulated, and transparent pathway to experimental treatments through licensed state institutions with board oversight.

76        (b) Condition payment on evidence development.

77        (c) Encourage coverage parity by state-regulated health  
78 plans.

79        (d) Align temporary pricing during the conditional period  
80 with the maturity of clinical evidence.

81        (3) As used in this section, the term:

82        (a) "Board" means the Board of Medicine, created under  
83 chapter 458, or the Board of Osteopathic Medicine, created under  
84 chapter 459, as applicable.

85        (b) "Department" means the Department of Health, created  
86 under s. 20.43.

87        (c) "Eligible facility" means a hospital or health care  
88 facility licensed under chapter 395 that operates under the  
89 federal policy for the protection of human subjects under 45  
90 C.F.R. part 46 and maintains, or participates under,  
91 Institutional Review Board oversight for experimental  
92 treatments.

93        (d) "Eligible patient" means a person who:

94        1.a. Has a terminal condition that is attested to by the  
95 treating physician and confirmed by a board-certified physician  
96 in an appropriate specialty for that condition; or

97        b. Has a life-threatening rare disease that is attested to  
98 by the treating physician and confirmed by a board-certified  
99 physician in an appropriate specialty.

100        2. Has considered and exhausted all other treatment

101 options for the terminal condition or life-threatening rare  
102 disease currently approved by the United States Food and Drug  
103 Administration.

104 3. Has given written informed consent pursuant to  
105 subsection (4) (c).

106 4. Has documentation from the treating physician that the  
107 patient meets the requirements of this paragraph.

108 (e) "Experimental treatment" means:

109 1. A drug, biological product, or medical device that has  
110 successfully completed phase 1 of a clinical trial but has not  
111 been approved for general use by the United States Food and Drug  
112 Administration (FDA) and remains under investigation in a  
113 clinical trial approved by the FDA; or

114 2. Personalized treatment for use by a patient based on  
115 his or her genomic or molecular profile, including, but not  
116 limited to, gene-targeted therapies, antisense oligonucleotides  
117 (ASOs)-mediated therapies, and neoantigen-targeting personalized  
118 cancer vaccines (PCVs), which have not been approved for general  
119 use by the FDA and remain under investigation in a clinical  
120 trial approved by the FDA.

121 (f) "Institutional Review Board" or "IRB" has the same  
122 meaning as in s. 381.86.

123 (g) "Patient registry" or "registry" means a clinical data  
124 collections system from which health information is collected at  
125 specific points during the course of treatment and all personal

126 identifiers have been removed or obscured to protect patient  
127 privacy.

128 (h) "Rare life-threatening disease" means a progressive  
129 disease or medical or surgical condition that affects less than  
130 200,000 persons in the United States. The term includes any  
131 progressive disease or medical or surgical condition that meets  
132 the criteria of a rare disease under 21 U.S.C. s. 360bb.

133 (i) "Terminal condition" means a progressive disease or  
134 medical or surgical condition that causes significant functional  
135 impairment, is not considered by a treating physician to be  
136 reversible even with the administration of available treatment  
137 options currently approved by the FDA, and, without the  
138 administration of life-sustaining procedures, will result in  
139 death within 1 year after diagnosis if the condition runs its  
140 normal course.

141 (4) (a) A physician licensed under chapter 458 or chapter  
142 459 may prescribe to an eligible patient an experimental  
143 treatment pursuant to this section.

144 (b)1. An eligible facility may administer to an eligible  
145 patient an experimental treatment if such treatment is  
146 administered pursuant to a platform or master protocol approved  
147 by an IRB pursuant to subsection (5) and complies with the  
148 registry requirements established in subsection (6).

149 2.a. An eligible facility that does not maintain its own  
150 IRB may establish and maintain an affiliation with another

151 eligible facility that maintains IRB oversight.

152 b. The department shall adopt rules for affiliations  
153 pursuant to this subparagraph, to include, but not be limited  
154 to, the handling of drugs, adverse-event escalation, quality  
155 assurance, and health data submission.

156 (c) Written informed consent, signed by an eligible  
157 patient, a court-appointed guardian for an eligible patient, or  
158 a health care surrogate designated by an eligible patient, is  
159 required, and must include:

160 1. An explanation of the currently approved treatments for  
161 the patient's terminal condition or life-threatening rare  
162 disease and that such treatments have been considered and  
163 exhausted.

164 2. Identification of the specific experimental treatment  
165 that the patient is seeking to use.

166 3. A description of the potential risks and benefits of  
167 using the experimental treatment, including best-case outcome  
168 and worst-case outcome. The description must be based on the  
169 physician's knowledge of the experimental treatment for the  
170 patient's terminal condition or life-threatening rare disease.

171 4. A statement that the patient's health plan or third-  
172 party administrator and physician are not obligated to pay for  
173 care or treatment consequent to the use of the experimental  
174 treatment unless required to do so by law or contract.

175 5. A statement that the patient's eligibility for hospice

176 care may be affected by the use of experimental treatment.

177 6. A statement that the patient understands he or she is  
178 liable for all expenses consequent to the use of the  
179 experimental treatment but that liability does not extend to the  
180 patient's estate, unless a contract between the patient and the  
181 manufacturer states otherwise.

182 7. An authorization from the patient to release his or her  
183 health information, collected at specific points during  
184 treatment, for submission into the registry established in  
185 subsection (6).

186 (5) A platform or master protocol must be approved by an  
187 IRB under which treating physicians and eligible facilities may  
188 adapt dosing, sequencing, or combinations of authorized  
189 experimental treatments, within predefined limitations, without  
190 seeking approval of an IRB for each adaptation. The platform or  
191 master protocol must specify safety limits, stopping  
192 requirements, and patient registry information pursuant to  
193 subsection (6).

194 (6) (a) The department shall establish and maintain a  
195 patient registry for the collection and analysis of experimental  
196 treatment outcomes under this section. Eligible facilities shall  
197 submit health information to the registry from which all  
198 personal identifiers have been removed or obscured to protect  
199 patient privacy.

200 (b) The department shall enter into a written contract



201 with eligible facilities which specifies the scope of services  
202 provided, the service level, the duration of the agreement, the  
203 responsible parties, and the service costs. Failure to meet the  
204 contract requirements may result in the filing of an action by  
205 the department and a temporary suspension of reimbursement for  
206 the prescription or administration of experimental treatment  
207 pursuant to subsection (8).

208 (c) The department shall annually publish a public report  
209 of experimental treatment outcomes and safety signals in the  
210 aggregate.

211 (7) (a) A health insurer or a health maintenance  
212 organization regulated under the Florida Insurance Code are  
213 encouraged to provide insurance coverage for experimental  
214 treatments in the same manner as any other covered treatment or  
215 therapy.

216 (b) A health insurer or a health maintenance organization  
217 may not deny insurance coverage based solely on a treatment  
218 being experimental or denied approval for general use by the  
219 FDA.

220 (c) This subsection does not require insurance coverage  
221 and does not expand the coverage an insurer or health  
222 maintenance organization must provide under the Florida  
223 Insurance Code.

224 (8) (a) A licensed physician or an eligible facility may  
225 receive reimbursement for the prescription or administration of

226 an experimental treatment under this section. However, in order  
227 to receive such reimbursement, an eligible facility must  
228 actively participate in the patient registry established under  
229 subsection (6). Failure to participate in the patient registry  
230 may result in temporary suspension of reimbursement until  
231 compliance is restored.

232 (b)1. Reimbursement for a prescribed or administered  
233 experimental treatment under this subsection may not exceed a  
234 discounted rate set by the department as a percentage of a  
235 recognized pricing benchmark, including wholesale acquisition  
236 cost or average sales price.

237 2. The department shall adopt rules that:

238 a. Establish an allowable discount band to reflect  
239 evidentiary uncertainty.

240 b. Apply a registry noncompliance adjustment which  
241 increases the discounted rate set by the department.

242 c. Provide for renewal and potential adjustment of the  
243 discounted rate every 2 years based on registry compliance.

244 (c) Upon approval by the FDA of an experimental treatment,  
245 the discounted rate set by the department for reimbursement  
246 expires and the standard reimbursement rate applies.

247 (9) A manufacturer of an experimental treatment shall  
248 provide for a temporary price concession, which price must be  
249 set by the department by rule as a percentage of a recognized  
250 pricing benchmark. The temporary price concession expires upon

251 approval by the FDA of the experimental treatment.

252 (10) An eligible facility or manufacturer and the  
253 Department of Management Services pursuant to s. 110.123 may  
254 enter into a healthcare contract under which a portion of  
255 payment is linked to experimental treatment outcomes. The  
256 department shall adopt a rule for such contract, which rule must  
257 provide for standard contract terms and authorize reconciliation  
258 intervals not to exceed 2 years.

259 (11) (a) The Department of Health shall adopt by rule  
260 procedures for licensing experimental treatment centers and  
261 shall approve or deny a completed application within 90 days.

262 (b) The department shall establish by rule the following  
263 minimum procedural and operational standards:

- 264 1. Written policies and procedures.
- 265 2. Oversight and governance standards.
- 266 3. Inspection and safety standards.
- 267 4. Staff training provisions.
- 268 5. Recordkeeping provisions.
- 269 6. Data quality assurance.

270 (12) (a) A licensing board may not revoke, deny renewal,  
271 suspend, or take any action against a physician's license issued  
272 under chapter 458 or chapter 459 based solely on the physician's  
273 recommendations to an eligible patient regarding access to or  
274 treatment with experimental treatment authorized under this  
275 section. A state agency responsible for Medicare certification

276 may not take action against a physician's Medicare certification  
277 based solely on the physician's recommendation that an eligible  
278 patient have access to experimental treatment.

279 (b)1. This section does not create a private cause of  
280 action:

281 a. Against a manufacturer of an experimental treatment;

282 b. Against a person or an entity involved in the care of  
283 an eligible patient who is using an experimental treatment; or

284 c. For any harm to the eligible patient that is a result  
285 of the use of an experimental treatment,

286  
287 if the manufacturer or other person or entity complies in good  
288 faith with the terms of this section and exercises reasonable  
289 care.

290 2. This paragraph does not apply to judicial action  
291 brought for gross negligence or intentional, willful, or wanton  
292 misconduct.

293 (c) If an eligible patient dies while using experimental  
294 treatment pursuant to this section, the patient's heirs are not  
295 liable for any outstanding debt related to the patient's use of  
296 such treatment.

297 (d) A hospital or licensed health care facility is not  
298 required to provide new or additional health care services or  
299 treatments under this section without prior consent of the  
300 hospital or licensed health care facility. Participation

301 pursuant to this section is entirely voluntary at all times.

302 (e) A public official, a public employee, or an agent of  
303 this state, or any political subdivision thereof, may not deny  
304 or attempt to deny access to experimental treatment under this  
305 section. Counseling, advice, or recommendations by a licensed  
306 physician consistent with the standard of care do not constitute  
307 the denial of access or the attempted denial of access to  
308 experimental treatment under this paragraph.

309 (13) This section does not authorize the possession, use,  
310 dispensing, or administration of a controlled substance in  
311 violation of chapter 893, except to the extent expressly  
312 permitted by state or federal law.

313 (14) The department shall adopt rules pursuant to ss.  
314 120.536(1) and 120.54 to implement this section.

315 **Section 3.** This act shall take effect upon becoming a law.