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SENATE BILL 1016

Health & Human Resources Committee Substitute Adopted 4/30/03

Short Title: Nursing Home/Medication Errors.	(Public)
Sponsors:	
Referred to:	
April 3, 2003	
A BILL TO BE ENTITLED	

1		A BILL TO BE ENTITLED
2	AN ACT R	EQUIRING NURSING HOMES TO ESTABLISH A MEDICATION
3	MANAGE	EMENT ADVISORY COMMITTEE AND SPECIFYING THE DUTIES
4	OF THE	COMMITTEE, AND TO REQUIRE NURSING HOMES TO DO
5	CERTAIN	THINGS PERTAINING TO THE REDUCTION OF
6	MEDICA	ΓΙΟΝ-RELATED ERRORS TO INCREASE PATIENT SAFETY.
7	The General A	Assembly of North Carolina enacts:
8	SE	CTION 1. Part 2 of Article 6 of Chapter 131E of the General Statutes is
9	amended by a	dding the following new sections to read:
10	" <u>§ 131E-128.</u>	1. Nursing home medication management advisory committee.
11	<u>(a)</u> Def	initions As used in this section, unless the context requires otherwise,
12	the term:	
13	<u>(1)</u>	'Advisory committee' means a medication management committee
14		established under this section to advise the quality assurance
15		<u>committee.</u>
16	<u>(2)</u>	'Medication-related error' means any preventable medication-related
17		event that adversely affects a patient in a nursing home and that is
18		related to professional practice, or health care products, procedures,
19		and systems, including, but not limited to, prescribing, prescription
20		order communications, product labeling, packaging and nomenclature,
21		compounding, dispensing, distribution, administration, education,
22		monitoring, and use.
23	<u>(3)</u>	'Nursing home' means a nursing home licensed under this Chapter and
24		includes an adult care home operated as part of a nursing home.
25	<u>(4)</u>	'Potential medication-related error' means a medication-related error
26		that has not yet adversely affected a patient in a nursing home, but that
27		has the potential to if not anticipated or prevented or if left unnoticed.
28	<u>(5)</u>	'Quality assurance committee' means a quality assurance committee
29		established in accordance with federal and State regulations to identify

1	issues with respect to which quality assessment and assume
1	issues with respect to which quality assessment and assurance
2 3	activities are necessary and to develop and implement appropriate
	plans of action to correct deficiencies in quality of care.
4	(b) Purpose. – It is the purpose of the General Assembly to enhance compliance
5	with this Part through the establishment of medication management advisory
6	committees in nursing homes. The purpose of these committees is to assist nursing
7	homes in identifying medication-related errors, evaluate the causes of those errors, and
8	take appropriate actions to ensure the safe prescribing, dispensing, and administration of
9	<u>medications to nursing home patients.</u>
10	(c) Advisory Committee Established; Membership. – Every nursing home shall
11	establish a medication management advisory committee to advise the quality assurance
12	committee on quality of care issues related to pharmaceutical and medication
13	management and use in the nursing home. The nursing home shall maintain the advisory
14	committee as part of its administrative duties. The advisory committee shall be
15	interdisciplinary and consist of the nursing home administrator and at least the
16 17	following members appointed by the nursing home administrator:
17	$(1) \qquad \frac{\text{The director of nursing.}}{\text{The consultant absence size}}$
18	(2) <u>The consultant pharmacist.</u> (2) <u>A physician designated by the pursing home administrator</u>
19 20	(3) <u>A physician designated by the nursing home administrator.</u>
20	(d) <u>At least three other members of the nursing home staff.</u>
21 22	(d) <u>Meetings. – The advisory committee shall meet as needed but not less</u>
	frequently than quarterly. The Director of Nursing or Staff Development Coordinator
23 24	shall chair the advisory committee. The nursing home administrator shall ensure that a
24 25	<u>record is maintained of each meeting.</u> (e) <u>Confidentiality. – The meetings or proceedings of the advisory committee,</u>
23 26	the records and materials it produces and the materials it considers, including analyses
20 27	and reports pertaining to medication-related error reporting under G.S. 131E-128.2 and
27	<u>G.S. 131E-128.5 and pharmacy reports on drug defects and adverse reactions under</u>
28 29	G.S. 131E-128.4, shall be confidential and not considered public records within the
29 30	meaning of G.S. 132-1, "Public records defined". The meetings or proceedings and
31	records and materials also shall not be subject to discovery or introduction into evidence
32	in any civil action against a nursing home licensed under this Article, or a provider of
33	professional health services which results from matters which are the subject of
34	evaluation and review by the committee. No person who was in attendance at a meeting
35	of the committee may testify in any civil action as to any evidence or other matters
36	produced or presented during the meetings or proceedings of the committee or as to any
37	findings, recommendations, evaluations, opinions, or other actions of the committee or
38	its members. Notwithstanding the foregoing:
39	(1) Information, documents, or records otherwise available, including any
40	deficiencies found in the course of an inspection conducted under G.S.
41	131E-105, shall not be immune from discovery or used in a civil action
42	merely because they were presented during meetings or proceedings of
43	the advisory committee. A member of the advisory committee or a
44	person who testifies before the committee may testify in a civil action
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1			but cannot be asked about that person's testimony before the
2			· ·
			committee or any opinion formed as a result of the committee
3		(0)	meetings or proceedings.
4		<u>(2)</u>	Information that is confidential and is not subject to discovery or use
5			in civil actions under this subsection may be released to a professional
6			standards review organization that performs any accreditation or
7			certification function. Information released to the professional
8			standards review organization shall be limited to that which is
9			reasonably necessary and relevant to the standards review
10			organizations' determination to grant or continue accreditation or
11			certification. Information released to the standards review organization
12			retains its confidentiality and is not subject to discovery or use in any
13			civil action as provided under this subsection, and the standards review
14			organization shall keep the information confidential subject to this
15			subsection.
16		(3)	Information that is confidential and is not subject to discovery or use
17		<u> </u>	in civil actions under this subsection may be released to the
18			Department of Health and Human Services pursuant to its investigative
19			authority under G.S. 131E-105. Information released to the
20			Department shall be limited to that which is reasonably necessary and
21			relevant to the Department's investigation of compliance with Part 1 of
22			Article 6 of this Chapter. Information released to the Department
23			retains its confidentiality and is not subject to discovery or use in any
24			civil action as provided in this subsection, and the Department shall
25			keep the information confidential subject to this subsection.
26		(4)	Information that is confidential and is not subject to discovery or use
27		<u>, , , , , , , , , , , , , , , , , , , </u>	in civil actions under this subsection may be released to an
28			occupational licensing board having jurisdiction over the license of an
29			individual involved in an incident that is under review or investigation
30			by the advisory committee. Information released to the occupational
31			licensing board shall be limited to that which is reasonably necessary
32			and relevant to an investigation being conducted by the licensing board
33			pertaining to the individual's involvement in the incident under review
34			by the advisory committee. Information released to an occupational
35			licensing board retains its confidentiality and is not subject to
36			discovery or use in any civil action as provided in this subsection, and
30 37			the occupational licensing board shall keep the information
38			confidential subject to this subsection.
30 39	(\mathbf{f})	Dutio	s. – The advisory committee shall do the following:
39 40	<u>(f)</u>		<u>Assess the nursing home's pharmaceutical management system,</u>
40 41		<u>(1)</u>	
41 42			including prescribing, distribution, administration policies, procedures,
42 43			and practices and identify areas at high risk for medication-related
43			errors.

1	(2)	Review the nursing home's pharmaceutical management goals and
2	<u>(2)</u>	respond accordingly to ensure that these goals are being met.
3	<u>(3)</u>	Review, investigate, and respond to nursing home incident reports,
4	<u>(5)</u>	deficiencies cited by licensing or credentialing agencies, and resident
5		grievances that involve actual or potential medication-related errors.
6	(A)	Identify goals and recommendations for the implementation of best
7	<u>(4)</u>	practices and procedures, including risk reduction technology, to
8		improve patient safety by reducing the risk of medication-related
8 9		errors.
10	(5)	Develop recommendations for the establishment of a mandatory,
11	<u>(0)</u>	nonpunitive, confidential reporting system within the nursing home of
12		actual and potential medication-related errors.
13	<u>(6)</u>	Develop specifications for drug dispensing and administration
14	<u>(0)</u>	documentation procedures to ensure compliance with federal and State
15		law, including the North Carolina Nursing Practice Act.
16	<u>(7)</u>	Develop specifications for self-administration of drugs by qualified
17	<u></u>	patients in accordance with law, including recommendations for
18		assessment procedures as to which patients may be qualified to
19		self-administer their medications.
20	(g) Penal	ty. – The Department may take adverse action against the license of a
21	-	upon a finding that the nursing home has failed to comply with this
22	-	1E-128.2, 131E-128.3, 131E-128.4, or 131E-128.5.
23	" <u>§ 131E-128.2</u>	. Nursing home quality assurance committee; duties related to
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24	(a) <u>medi</u> (a) <u>Every</u> assurance comr	<u>Nursing home quality assurance committee; duties related to</u> <u>cation error prevention.</u> <u>nursing home administrator shall ensure that the nursing home quality</u> <u>nittee develops and implements appropriate measures to minimize the</u>
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1	(8)	Implement policies and procedures designed to improve accuracy in
2	<u>(0)</u>	medication administration and in documentation by properly
3		authorized individuals, in accordance with prescribed orders and stop
4		order policies.
4 5	(9)	
5 6	<u>(9)</u>	<u>Implement policies and procedures for the self-administration of</u> medication.
0 7	(10)	
	<u>(10)</u>	Investigate and analyze the frequency and root causes of general
8 9		categories and specific types of actual or potential medication-related
9 10	(11)	errors. Develop recommendations for plans of action to correct identified
	<u>(11)</u>	Develop recommendations for plans of action to correct identified
11	"8 121E 190 2	deficiencies in the facility's pharmaceutical management practices.
12		Staff orientation on medication error prevention.
13	-	home administrator shall ensure that the nursing home provide a
14		e hour of education and training in the prevention of actual or potential
15		ted errors. This training shall be provided upon orientation and annually
16		nonphysician personnel involved in direct patient care. The content of
17	-	l include at least the following:
18	<u>(1)</u>	General information relevant to the administration of medications
19		including terminology, procedures, and routes of medication
20		administration, potential side effects and adverse reactions.
21	<u>(2)</u>	Additional instruction on categories of medication pertaining to the
22		specific needs of the patient receiving the medication.
23	<u>(3)</u>	The facility's policy and procedures regarding its medication
24		administration system.
25	<u>(4)</u>	How to assist patients with safe and accurate self-administration,
26		where appropriate.
27	<u>(5)</u>	Identifying and reporting actual and potential medication-related
28		errors.
29	" <u>§ 131E-128.4.</u>	Nursing home pharmacy reports; duties of consultant pharmacist.
30	$\underline{(a)}$ The c	onsultant pharmacist shall conduct a drug regimen review for actual and
31	*	therapy problems and make remedial or preventive clinical
32	recommendation	ns into the medical record of every patient receiving medication. The
33	consultant pharm	macist shall conduct the review at least monthly in accordance with the
34	nursing home's	policies and procedures.
35	<u>(b)</u> <u>The c</u>	consultant pharmacist shall report and document any drug irregularities
36	and clinical rec	ommendations promptly to the attending physician or nurse-in-charge
37	and the nursing	home administrator. The reports shall include problems identified and
38	recommendation	ns concerning:
39	(1)	Drug therapy which may be affected by biological agents, laboratory
40		tests, special dietary requirements, and foods used or administered
41		concomitantly with other medication to the same recipient.
42	(2)	Monitoring for potential adverse effects.
43	$\overline{(3)}$	Allergies.
	<u> </u>	U

1	(4) Drug interactions, including interactions between prescription drugs
2	and over-the-counter drugs, drugs and disease, and interactions
3	between drugs and nutrients.
4	(5) <u>Contraindications and precautions.</u>
5	(6) <u>Potential therapeutic duplication.</u>
6	(7) Over-extended length of treatment of certain drugs typically prescribed
7	for a short period of time.
8	(8) Beer's listed drugs which are potentially inappropriate for use by
9	elderly persons.
10	(9) <u>Under treatment or medical conditions that are suboptimally treated or</u>
11	not treated at all that warrant additional drug therapy to ensure quality
12	<u>of care.</u>
13	(10) Other identified problems and recommendations.
14	(c) The consultant pharmacist shall report drug product defects and adverse drug
15	reactions in accordance with the ASHSP-USP-FDA Drug Product Defect Reporting
16	System and the USP Adverse Drug Reaction Reporting System. The term "ASHSP-
17	USP-FDA" means American Society of Health System Pharmacists-United States
18	Pharmacopoeia-Food and Drug Administration. Information released to the ASHSP-
19	USP-FDA retains its confidentiality and is not subject to discovery or use in any civil
20	actions as provided under G.S. 131E-128.1.
21	(d) The consultant pharmacist shall ensure that all known allergies and adverse
22	effects are documented in plain view in the patient's medical record including the
23	medication administration records and communicated to the dispensing pharmacy. The
24	specific medications and the type of allergy or adverse reaction shall be specified in the
25	documentation.
26	(e) The consultant pharmacist shall ensure that drugs that are not specifically
27	limited as to duration of use or number of doses shall be controlled by automatic stop
28	orders. The pharmacist consultant shall further ensure that the prescribing provider is
29	notified of the automatic stop order prior to the dispensing of the last dose so that the
30	provider may decide whether to continue to use the drug.
31	(f) The consultant pharmacist shall, on a quarterly basis, submit a summary of
32	the reports submitted under subsections (a) and (b) of this section to the medication
33	management advisory committee established under G.S. 131E-128.1. The summary
34	shall not include any information that would identify a patient, a family member, or an
35	employee of the nursing home. The purpose of the summary shall be to facilitate the
36	identification and analysis of weaknesses in the nursing home's pharmaceutical care
37	system that have an adverse impact on patient safety.
38	"§ 131E-128.5. Medication-related error reports.
39	(a) The Secretary of Health and Human Services shall contract with a public or
40	private entity to develop and implement a Medication Error Quality Initiative. The
41	Initiative would provide for, among other things, receipt and analysis by the contracting
42	entity of annual reports from each nursing home on the nursing home's medication-
43	related errors. The report submitted by the nursing home shall not contain information

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1	that would iden	tify the patient, individual reporting the error, or other persons involved
2	in the occurrence	e. The report shall include the following:
3	<u>(1)</u>	The total number of medication-related errors for the preceding year.
4	<u>(2)</u>	A listing of the types of medication-related errors, the number of
5		medication-related errors, the root cause analysis of each error, and the
6		staff level involved.
7	<u>(3)</u>	A listing of the types of injury caused and the number of injuries
8		occurring.
9	<u>(4)</u>	Types of liability claims filed based on an adverse incident or
10		<u>reportable injury.</u>
11	<u>(b)</u> The	contracting entity shall provide for analysis of the medication-related
12	error reports to	determine trends in the incidence of medication-related errors in nursing
13	homes. Inform	ation released to the contractor retains its confidentiality and is not
14	subject to disco	very or use in any civil actions as provided under G.S. 131E-128.1, and
15	the contractor s	hall keep the information confidential subject to that section."
16	SEC'	FION 2. The Department shall use available grants and federal funds to
17	implement G.S.	131E-182.5 as enacted in this act.
18	SEC	FION 3. This act becomes effective January 1, 2004.