1	10A NCAC 13D .2306 is proposed for amendment as follows:		
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3	10A NCAC 13D .2306 MEDICATION ADMINISTRATION		
4	(a) The facility shall ensure that medications are administered in accordance with standards of professional practice		
5	and applicable occupational licensure regulations. regulations and manufacturer's recommendations.		
6	(b) The facility shall ensure that each patient's drug regimen is free from drugs used in excessive dose or duplicative		
7	therapy, for excessive duration or without adequate indications for the prescription of the drug. Drugs shall not be		
8	used without adequate monitoring or in the presence of adverse conditions that indicate the drugs' usage should be		
9	modified or discontinued. As used in this paragraph:		
10	(1) "Excessive dose" means the total amount of any medication (including duplicate therapy) given at		
11	one time or over a period of time that is greater than the amount recommended by the		
12	manufacturer for a resident's age and condition.		
13	(2) "Excessive Duration" means the medication is administered beyond the manufacturer's		
14	recommended time frames or facility-established stop order policies or without either evidence of		
15	additional therapeutic benefit for the resident or clinical evidence that would warrant the continued		
16	use of the medication.		
17	(3) "Duplicative Therapy" means multiple medications of the same pharmacological class or category		
18	or any medication therapy that replicates a particular effect of another medication that the		
19	individual is taking.		
20	(4) "Indications for the prescription" means a documented clinical rationale for administering a		
21	medication that is based upon an assessment of the resident's condition and therapeutic goals and		
22	is consistent with manufacturer's recommendations.		
23	(5) "Monitoring" means ongoing collection and analysis of information (such as observations and		
24	diagnostic test results) and comparison to baseline data in order to:		
25	(A) Ascertain the individual's response to treatment and care, including progress or lack of		
26	progress toward a therapeutic goal;		
27	(B) Detect any complications or adverse consequences of the condition or of the treatments		
28	and;		
29	(C) Support decisions about modifying, discontinuing, or continuing any interventions.		
30	(c) Antipsychotic therapy shall not be initiated on any patient unless necessary to treat a clinically diagnosed and		
31	clinically documented condition. When antipsychotic therapy is prescribed, unless clinically contraindicated,		
32	gradual dose reductions and behavioral interventions shall be employed in an effort to discontinue these drugs.		
33	"Gradual dose reduction" means the stepwise tapering of a dose to determine if symptoms, conditions or risks can be		
34	managed by a lower dose or if the dose or the medication can be discontinued.		

(d) The facility shall ensure that procedures aimed at minimizing medication error rates include, but are not limited

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to, include the following:

1	(1)	All medications or drugs and treatments shall be administered and discontinued in accordance
2		with signed medical orders which are recorded in the patient's medical record. Such orders shall be
3		complete and include drug name, strength, quantity to be administered, route of administration,
4		frequency and, if ordered on an as-needed basis, a elearly stated indication for use.
5	(2)	The requirements for self-administration of medication shall include, but not be limited to, include
6		the following:
7		(A) determination by the interdisciplinary team that this practice is safe;
8		(B) administration ordered by the physician or other person legally authorized to prescribe
9		medications;
10		(C) specific instructions for administration printed on the medication label; and
11		(D) administration of medication monitored by the licensed nursing staff and consultant
12		pharmacist.
13	(3)	The administration of one patient's medications to another patient is prohibited except in the case
14		of an emergency. In the event of such emergency, steps shall be taken to the facility shall ensure
15		that the borrowed medications are replaced promptly and so documented.
16	(4)	Omission of medications and the reason for omission shall be indicated in the patient's medical
17		record.
18	(5)	Medication administration records shall provide time of administration, identification of the drug
19		and strength of drug, quantity of drug administered, route of administration, frequency,
20		documentation sufficient to determine the staff who administered the drugs. Medication
21		administration records shall indicate documentation of injection sites and topical medication sites
22		requiring rotation, including, but not limited to, rotation of transdermal medication.
23	(6)	The pharmacy shall receive an exact copy of each physician's order for medications and
24		treatments.
25	(7)	When medication orders do not state the number of doses or days to administer the medication, the
26		facility shall implement automatic Automatic stop orders for medications and treatments shall be
27		established and implemented. according to manufacturer's recommendations.
28	(8)	The facility shall maintain an accountability of controlled substances as defined by the North
29		Carolina Controlled Substances Act, G.S. 90, Article 5.
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31	History Note:	Authority G.S. 131E-104;
32		Eff. January 1, 1996. <u>1996;</u>
33		Amended Eff. January 1, 2013.