## GENERAL ASSEMBLY OF NORTH CAROLINA SESSION 2025

H D

## HOUSE BILL 330 PROPOSED COMMITTEE SUBSTITUTE H330-PCS30211-CV-9

Short Title:	Controlled Substances Act - Updates.		
Sponsors:			
Referred to:			

## March 10, 2025

	Watch 10, 2023			
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2	AN ACT TO UPDATE THE CONTROLLED SUBSTANCES ACT.			
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22	dddd. Phenyl U-47700.			
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                              N,N-didesmethyl U-47700.
                       SSSS.
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                              U-62066.
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                       uuuu. Propyl U-47700.
 4
                       vvvv. (2,3- or 3,4)-Ethylenedioxy U-51754.
 5
                                  4-phenyl U-51754.
                       wwww.
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                       xxxx. N-desmethyl U-47700.
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                       yyyy. (2,3- or 3,4)-Ethylenedioxy U-47700.
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                       zzzz. N-methyl U-47931E.
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                       aaaaa. (2,3- or 3,4)-Methylenedioxy U-47700.
10
                       bbbbb. U-69593.
11
                       ccccc. U-50488.
                       ddddd. U-48753E.
12
                       eeeee. U-47931E."
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                SECTION 1.(b) G.S. 90-89(1a) reads as rewritten:
14
                "(1a) Fentanyl derivatives. – Unless specifically excepted, listed in another
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                       schedule, or contained within a pharmaceutical product approved by the
16
                       United States Food and Drug Administration, any compound structurally
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18
                       derived from N-[1-(2-phenylethyl)-4-piperidinyl]-N-phenylpropanamide
19
                       (Fentanyl) by any substitution on or replacement of the phenethyl group, any
20
                       substitution on the piperidine ring, any substitution on or replacement of the
21
                       propanamide group, any substitution on the anilido phenyl group, or any
                       combination of the above unless specifically excepted or listed in another
22
                       schedule to include their salts, isomers, and salts of isomers. Fentanyl
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24
                       derivatives include, but are not limited to, the following:
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26
                       f.
27
                                  N-(2-fluorophenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-propana
28
                                      (also known as 2-fluorofentanyl).(also known as
29
                              ortho-fluorofentanyl).
30
                       g.
31
                                  N-(3-fluorophenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-propana
32
                                      (also known as 3-fluorofentanyl).(also known as
33
                              meta-fluorofentanyl).
34
                       . . .
35
                       i.
36
                                  N-(4-fluorophenyl)-2-methyl-N-[1-(2-phenylethyl)-4-piperidinyl]
37
                              -propanamide (also known as 4-fluoroisobutyryl fentanyl,
                              4-FIBF): (also known as 4-fluoroisobutyryl fentanyl).
38
39
                              N-(4-fluorophenyl)-N-[1-(2-phenylethyl)-4-piperidinyl]-butanamide
                       j.
40
                              (also known as 4-fluorobutyryl fentanyl, 4-FBF).(also known as
41
                              para-fluorobutyryl fentanyl).'
42
                SECTION 1.(c) G.S. 90-89 is amended by adding a new subdivision to read:
43
                "(1b) Nitazene derivatives. – The N-substituted benzimidazole structural class,
                       including any of the following derivatives, their salts, isomers, or salts of
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                       isomers unless specifically utilized as part of the manufacturing process by a
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                       commercial industry of a substance or material not intended for human
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                       ingestion or consumption, as a prescription administered under medical
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                       supervision, or for research at a recognized institution, whenever the existence
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                       of these salts, isomers, or salts of isomers is possible within the specific
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                       chemical designation or unless specifically excepted or listed in this or another
                       schedule, structurally derived from benzimidazole by substitution at the
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1-position nitrogen with an ethylamine group, and by substitution at the 1 2 2-position carbon with a benzyl group, whether or not the compound is further 3 modified in any of the following ways: 4 By monoalkyl or dialkyl substitution on the 1'-nitrogen of the 5 1-position ethylamine group, or by inclusion of the nitrogen in a cyclic 6 7 By substitution on the 2'-methylene carbon of the benzyl group by <u>b.</u> 8 alkyl or carboxamide groups. 9 By replacement of the 2'-methylene carbon group with an ethylbenzyl, <u>c.</u> thiophenol, or methoxybenzene group, which may be further 10 11 substituted with alkyl, hydroxyl, alkoxy, acetoxy, halide, or sulfide 12 groups. 13 By substitution at the 2'-position, 3'-position, or 4'-position of the d. 14 benzyl group, or both, with alkyl, hydroxyl, alkoxy, acetoxy, halide, or sulfide groups. 15 By replacement of a phenyl hydrogen atom at either the 5-position or 16 <u>e.</u> 17 6-position of the benzimidazole core with a nitro, or primary amine 18 group." **SECTION 1.(d)** G.S. 90-89(3)mm. reads as rewritten: 19 20 "mm. 5-methoxy N-methyl N-propyltryptamine (5 MeO MiPT).5-methoxy-N-methyl-N-isopropyltryptamine 21 22 (5-MeO-MiPT)." **SECTION 1.(e)** G.S. 90-89(4) is amended by adding a new sub-subdivision to read: 23 24 Bromazolam." 25 **SECTION 1.(f)** G.S. 90-89(5)j. reads as rewritten: 26 "į. Substituted cathinones. A compound, other than bupropion, that is 27 derived from 2-amino-1-phenyl-1-propanone 28 modification in any of the following ways: (i) by substitution in the 29 phenyl ring to any extent with alkyl, alkoxy, alkylenedioxy, haloalkyl, 30 or halide substituents, whether or not further substituted in the phenyl 31 ring by one or more other univalent substituents; (ii) by substitution at 32 the 3-position to any extent; or (iii) by substitution at the nitrogen atom 33 with alkyl, dialkyl, benzyl, cycloalkyl, or methoxybenzyl groups or by 34 inclusion of the nitrogen atom in a cyclic structure. For the purpose of 35 this paragraph, the term "isomer" includes the optical, positional, or 36 geometric isomer." **SECTION 1.(g)** G.S. 90-89(7) reads as rewritten: 37 Synthetic cannabinoids. – Any quantity of any synthetic chemical compound 38 "(7)39 that (i) is a cannabinoid receptor agonist and mimics the pharmacological 40 effect of naturally occurring substances or (ii) has a stimulant, depressant, or 41 hallucinogenic effect on the central nervous system that is not listed as a 42 controlled substance in Schedules I through V, and is not an FDA-approved 43 drug. Synthetic cannabinoids include, but are not limited to, the substances listed in sub-subdivisions a. through p. v. of this subdivision and any substance 44 45 that contains any quantity of their salts, isomers (whether optical, positional, 46 or geometric), homologues, and salts of isomers and homologues, unless specifically excepted, whenever the existence of these salts, isomers, 47 homologues, and salts of isomers and homologues is possible within the 48 49 specific chemical designation. The following substances are examples of

synthetic cannabinoids and are not intended to be inclusive of the substances

included in this Schedule:

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*l*. Indole carboxamides. Any compound structurally derived from 1H-indole-3-carboxamide or 1H-indole-2-carboxamide substituted in one or both of the following ways:

- 1. At the nitrogen atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-3-morpholinyl)methyl, tetrahydropyranylmethyl, benzyl, or halo benzyl group; andor
- 2. At the nitrogen of the carboxamide by a phenyl, benzyl, naphthyl, adamantyl, cyclopropyl, or propionaldehyde group;group, or methyl 3,3-dimethyl-butanoate group; whether or not the compound is further modified to any extent in the following ways: (i) substitution to the indole ring to any extent, (ii) substitution to the phenyl, benzyl, naphthyl, adamantyl, cyclopropyl, or propionaldehyde group to any extent, (iii) a nitrogen heterocyclic analog of the indole ring, or (iv) a nitrogen heterocyclic analog of the phenyl, benzyl, naphthyl, adamantyl, or cyclopropyl ring. Substances in this class include, but are not limited to: SDB-001 and STS-135.STS-135 and MDMB-ICA.

n. Indazole carboxaldehydes. Any compound structurally derived from 1H-indazole-3-carboxaldehyde or 1H-indazole-2-carboxaldehyde substituted in both of the following ways:

2. At the carbon of the carboxaldehyde by a phenyl, benzyl, naphthyl, adamantyl, cyclopropyl, or propionaldehyde group; whether or not the compound is further modified to any extent in the following ways: (i) substitution to the indazole ring to any extent, (ii) substitution to the phenyl, benzyl, naphthyl, adamantyl, cyclopropyl, or propionaldehyde group to any extent, (iii) a nitrogen heterocyclic analog of the indazole ring, or (iv) a nitrogen heterocyclic analog of the phenyl, benzyl, naphthyl, adamantyl, or cyclopropyl ring.

- o. Indazole carboxamides. Any compound structurally derived from 1H-indazole-3-carboxamide or 1H-indazole-2-carboxamide substituted in <u>one or</u> both of the following ways:
  - 1. At the nitrogen atom of the indazole ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-3-morpholinyl)methyl, tetrahydropyranylmethyl, benzyl, or halo benzyl group; andor
  - 2. At the nitrogen of the carboxamide by a phenyl, benzyl, naphthyl, adamantyl, cyclopropyl, or propionaldehyde group; group, or methyl 3,3-dimethyl-butanoate group;

whether or not the compound is further modified to any extent in the following ways: (i) substitution to the indazole ring to any extent, (ii) substitution to the phenyl, benzyl, naphthyl, adamantyl, cyclopropyl,

1		or propionaldehyde group to any extent, (iii) a nitrogen heterocyclic
2		analog of the indazole ring, or (iv) a nitrogen heterocyclic analog of
3		the phenyl, benzyl, naphthyl, adamantyl, or cyclopropyl ring.
4		Substances in this class include, but are not limited to: AKB-48,
5		fluoro-AKB-48, APINCACA, AB-PINACA, AB-FUBINACA,
6		ADB-FUBINACA, and ADB-PINACA. ADB-PINACA.
7		ADB-INACA, MDMB-INACA, MDMB-5Me-INACA, and
8		MDMB-5Br-INACA.
9		
10	<u>s.</u>	Oxindoles. Any compound structurally derived from
11		3-hydrazonoindolin-2-one substituted in one or both of the following
12		ways:
13		1. At the nitrogen atom of the oxoindole ring by an alkyl,
14		haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl,
15		cycloalkylethyl; or
16		2. At the nitrogen of the hydrazide by a phenyl, benzyl, naphthyl,
17		adamantyl, cyclopropyl, or propionaldehyde group;
18		whether or not the compound is further modified to any extent
19		in the following ways: (i) substitution to the oxoindole ring to
20		any extent or (ii) substitution to the phenyl, benzyl, naphthyl,
21		adamantyl, cyclopropyl, or propionaldehyde group to any
22		extent. Substances in this class include, but are not limited to:
23		BZO-POXIZID, BZO-HEXOXIZIDE, 5F-BZO-POXIZIDE.
24	<u>t.</u>	Indole acetamides. Any compound structurally derived from
25	<u>u.</u>	1H-indole-3-acetamide or 1H-indole-2-acetamide substituted in one or
26		both of the following ways:
27		1. At the nitrogen atom of the indole ring by an alkyl, haloalkyl,
28		cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl,
29		1-(N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl,
30		1-(N-methyl-2-pyrrolidinyl)methyl,
31		1-(N-methyl-3-morpholinyl)methyl, tetrahydropyranylmethyl,
32		benzyl, or halo benzyl group; or
33		- · · · · · · · · · · · · · · · · · · ·
		2. At the nitrogen of the acetamide by a phenyl, benzyl, naphthyl, adamantyl, cyclopropyl, or propionaldehyde group;
34 35		- · · · · · · · · · · · · · · · · · · ·
		whether or not the compound is further modified to any extent
36		in the following ways: (i) substitution to the indole ring to any
37		extent, (ii) substitution to the phenyl, benzyl, naphthyl,
38		adamantyl, cyclopropyl, or propionaldehyde group to any
39		extent, (iii) a nitrogen heterocyclic analog of the indole ring, or
40		(iv) a nitrogen heterocyclic analog of the phenyl, benzyl,
41		naphthyl, adamantyl, or cyclopropyl ring. Substances in this
42		class include, but are not limited to: AFUBIATA, CH-PIATA
43		AB-CHMIATA, ADB-FUBIATA.
44	<u>u.</u>	Indazole acetaldehydes. Any compound structurally derived from
45		1H-indazol-3-ylacetaldehyde or 1H-indazol-2-ylacetaldehyde
46		substituted in one or both of the following ways:
47		1. At the nitrogen atom of the indazole ring by an alkyl, haloalkyl,
48		cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl,
49		1-(N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl,
50		1-(N-methyl-2-pyrrolidinyl)methyl,

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1	1-(N	I-methyl-3-morpholinyl)methyl, tetrahydropyranylmethyl,
2		zyl, or halo benzyl group; or
3		the nitrogen of the carboxamide by a phenyl, benzyl,
4		nthyl, adamantyl, cyclopropyl, or propionaldehyde group;
5	whe	ther or not the compound is further modified to any extent
6	<u>in tl</u>	ne following ways: (i) substitution to the indazole ring to
7		extent, (ii) substitution to the phenyl, benzyl, naphthyl,
8		nantyl, cyclopropyl, or propionaldehyde group to any
9	exte	nt, (iii) a nitrogen heterocyclic analog of the indazole ring,
10	or (	iv) a nitrogen heterocyclic analog of the phenyl, benzyl,
11		nthyl, adamantyl, or cyclopropyl ring. Substances in this
12		s include, but are not limited to: ADB-BUTINAATA,
13		B-FUBINAATA.
14		Any compound structurally derived from 1H-pyrazole
15		in all of the following ways:
16		he 1 position of the pyrazole ring by an alkyl, haloalkyl, or
17		<u>nyl group.</u>
18		he 3 position of the pyrazole ring by a halo benzyl or
19		pionaldehyde group.
20		he 5 position of the pyrazole ring by a halo benzyl or
21	<del></del>	pionaldehyde group;
22		ther or not the compound is further modified by a
23		stitution to the propionaldehyde group to any extent.
24		stances in this class include, but are not limited to:
25		ADB-4en-PFUPPYCA, 5-fluoro-3,5-AB-PFUPPYCA."
26 27	` ,	90(2)h1. reads as rewritten: immediate precursor chemical,
28	<b>5</b>	1
28 29		-phenethyl-4-piperidine
30	SECTION 1.(i) G.S. 90-9	nilino-N-phenethylpiperdine (ANPP)."
31		yltestosterone, Dehydrochloromethyltestosterone,"
32	3	91(k)16. reads as rewritten:
33	"16. Mesterolene, Meste	
34	·	effective when it becomes law.
54	SECTION 2. This act is t	dictive when it decomes law.

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