23.8099.03000

Sixty-eighth Legislative Assembly of North Dakota

FIRST ENGROSSMENT with House Amendments ENGROSSED SENATE BILL NO. 2093

Introduced by

Judiciary Committee

(At the request of the State Board of Pharmacy)

- 1 A BILL for an Act to amend and reenact sections 19-03.1-05, 19-03.1-11, and 19-03.1-13 of the
- 2 North Dakota Century Code, relating to the scheduling of controlled substances; and to declare
- 3 an emergency.

4 BE IT ENACTED BY THE LEGISLATIVE ASSEMBLY OF NORTH DAKOTA:

- 5 **SECTION 1. AMENDMENT.** Section 19-03.1-05 of the North Dakota Century Code is
- 6 amended and reenacted as follows:
- 7 19-03.1-05. Schedule I.
- 8 1. The controlled substances listed in this section are included in schedule I.
- 9 2. Schedule I consists of the drugs and other substances, by whatever official name,
- 10 common or usual name, chemical name, or brand name designated, listed in this
- 11 section.
- 12 3. Opiates. Unless specifically excepted or unless listed in another schedule, any of the
- following opiates, including their isomers, esters, ethers, salts, and salts of isomers,
- esters, and ethers, whenever the existence of those isomers, esters, ethers, and salts
- is possible within the specific chemical designation:
- 16 a. Acetylmethadol.
- b. Allylprodine.
- 18 c. Alphacetylmethadol.
- d. Alphameprodine.
- e. Alphamethadol.
- 21 f. Benzethidine.
- g. Betacetylmethadol.
- h. Betameprodine.
- i. Betamethadol.

1	j.	Betaprodine.
2	k.	Brorphine.
3	l.	Clonitazene.
4	m.	Dextromoramide.
5	n.	Diampromide.
6	0.	Diethylthiambutene.
7	p.	Difenoxin.
8	q.	Dimenoxadol.
9	r.	Dimepheptanol.
10	S.	Dimethylthiambutene.
11	t.	Dioxaphetyl butyrate.
12	u.	Dipipanone.
13	V.	Ethylmethylthiambutene.
14	W.	Etonitazene.
15	Х.	Etoxeridine.
16	y.	Furethidine.
17	Z.	Hydroxypethidine.
18	aa.	Isotonitazene (also known as N,N-diethyl-2-(2-(4- isopropoxybenzyl)-5-nitro-1H-
19		benzimidazol-1-yl)ethan-1-amine).
20	bb.	Ketobemidone.
21	CC.	Levomoramide.
22	dd.	Levophenacylmorphan.
23	ee.	Morpheridine.
24	ff.	MPPP (also known as 1-methyl-4-phenyl-4-propionoxypiperidine).
25	99	Noracymethadol.
26	hh.	Norlevorphanol.
27	ii.	Normethadone.
28	jj.	Norpipanone.
29	kk.	PEPAP (1-(2-Phenylethyl)-4-Phenyl-4-acetoxypiperidine).
30	II.	Phenadoxone.
31	mm.	Phenampromide.

1	nn.	Phenomorphan.
2	00.	Phenoperidine.
3	pp.	Piritramide.
4	qq.	Proheptazine.
5	rr.	Properidine.
6	SS.	Propiram.
7	tt.	Racemoramide.
8	uu.	Tilidine.
9	VV.	Trimeperidine.
10	WW.	3,4-dichloro-N-[2-(dimethylamino)cyclohexyl]-N-methylbenzamide (also known as
11		U-47700).
12	XX.	1-cyclohexyl-4-(1,2-diphenylethyl)piperazine (also know as MT-45).
13	уу.	3,4-dichloro- <i>N</i> -{[1-(dimethylamino)cyclohexyl]methyl}benzamide (also known as
14		AH-7921).
15	ZZ.	Zipeprol.
16	aaa.	2-(2-(4-butoxybenzyl)-5-nitro-1H-benzimidazol-1-yl)-N,N-diethylethan-1-amine
17		(also known as Butonitazene).
18	<u>bbb.</u>	2-(2-(4-ethoxybenzyl)-1H-benzimidazol-1-yl)-N,N-diethylethan-1-amine (also
19		known as Etodesnitazene and etazene).
20	CCC.	N,N-diethyl-2-(2-(4-fluorobenzyl)-5-nitro-1H-benzimidazol-1-yl)ethan-1-amine
21		(also known as Flunitazene).
22	<u>ddd.</u>	N,N-diethyl-2-(2-(4-methoxybenzyl)-1H-benzimidazol-1-yl)ethan-1-amine (also
23		known as Metodesnitazene).
24	eee.	N,N-diethyl-2-(2-(4-methoxybenzyl)-5-nitro-1H-benzimidazol-1-yl)ethan-1-amine
25		(also known as Metonitazene).
26	<u>fff.</u>	2-(4-ethoxybenzyl)-5-nitro-1-(2-(pyrrolidin-1-yl)ethyl)-1H-benzimidazole (also
27		known as N-Pyrrolidino Etonitazene and Etonitazepyne).
28	<u>ggg.</u>	N,N-diethyl-2-(5-nitro-2-(4-propoxybenzyl)-1H-benzimidazol-1-yl)ethan-1-amine
29		(also known as Protonitazene).
30	<u>hhh.</u>	Fentanyl derivatives. Unless specifically excepted or unless listed in another
31		schedule or are not FDA approved drugs, and are derived from N-(1-(2-

1	Phe	Phenylethyl)-4-piperidinyl)-N-phenylpropanamide (Fentanyl) by any substitution					
2	on o	on or replacement of the phenethyl group, any substitution on the piperidine ring,					
3	any	any substitution on or replacement of the propanamide group, any substitution on					
4	the	anilido phenyl group, or any combination of the above. Examples include:					
5	(1)	N-[1-(1-methyl-2-phenethyl)-4-piperidinyl]-N-phenylacetamide (also known					
6		as Acetyl-alpha-methylfentanyl).					
7	(2)	N-[1-(alpha-methyl-beta-phenyl)ethyl-4-piperidyl]propionanilide; 1-(1-methyl-					
8		2-phenylethyl)-4-(N-propanilido)piperidine (also known as Alpha-					
9		methylfentanyl).					
10	(3)	N-[1-methyl-2-(2-thienyl)ethyl-4-piperidinyl]-N-phenylpropanamide (also					
11		known as Alpha-methylthiofentanyl).					
12	(4)	N-[1-(2-hydroxy-2-phenethyl)-4-piperidinyl]-N-phenylpropanamide (also					
13		known as Beta-hydroxyfentanyl).					
14	(5)	N-[1-(2-hydroxy-2-phenethyl)-3-methyl-4-piperidinyl]-N-phenylpropanamide					
15		(also known as Beta-hydroxy-3-methylfentanyl).					
16	(6)	N-[3-methyl-1-(2-phenylethyl)-4-piperidyl]-N-phenylpropanamide (also					
17		known as 3-Methylfentanyl).					
18	(7)	N-[3-methyl-1-(2-thienyl)ethyl-4-piperidinyl]-N-phenylpropanamide (also					
19		known as 3-Methylthiofentanyl).					
20	(8)	N-(4-fluorophenyl)-N-[1-(2-phenethyl)-4-piperidinyl]propanamide (also					
21		known as Para-fluorofentanyl).					
22	(9)	N-phenyl-N-[1-(2-thienyl)ethyl-4-piperidinyl]propanamide (also known as					
23		Thiofentanyl).					
24	(10)	N-(1-phenylethylpiperidin-4-yl)-N-phenylfuran-2-carboxamide (also known					
25		as Furanyl Fentanyl).					
26	(11)	N-(1-phenethylpiperidin-4-yl)-N-phenylbutyramide; N-(1-phenethylpiperidin-					
27		4-yl)-N-phenylbutanamide (also known as Butyryl Fentanyl).					
28	(12)	N-[1-[2-hydroxy-2-(thiophen-2-yl)ethyl]piperidin-4-yl]-N-phenylpropionamide;					
29		N-[1-[2-hydroxy-2-(2-thienyl)ethyl]-4-piperidinyl]-N-phenylpropanamide (also					
30		known as Beta-Hydroxythiofentanyl).					

1	(13)	N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide (also known as Acetyl
2		Fentanyl).
3	(14)	N-(1-phenethylpiperidin-4-yl)-N-phenylacrylamide (also known as Acryl
4		Fentanyl).
5	(15)	N-(1-phenethylpiperidin-4-yl)-N-phenylpentanamide (also known as Valeryl
6		Fentanyl).
7	(16)	N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide (also known
8		as 4-Fluoroisobutyryl Fentanyl).
9	(17)	N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)propionamide (also known
10		as Ortho-fluorofentanyl, 2-Fluorofentanyl).
11	(18)	N-(1-phenethylpiperidin-4-yl)-N-phenyltetrahydrofuran-2-carboxamide (also
12		known as Tetrahydrofuranyl Fentanyl).
13	(19)	2-methoxy-N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide (also known as
14		Methoxyacetyl Fentanyl).
15	(20)	N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopropanecarboxamide (also
16		known as Cyclopropyl Fentanyl).
17	(21)	N-(2-fluorophenyl)-2-methoxy-N-(1-phenethylpiperidin-4-yl)acetamide (also
18		known as Ocfentanil).
19	(22)	N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopentanecarboxamide (also
20		known as Cyclopentyl Fentanyl).
21	(23)	N-(1-phenethylpiperidin-4-yl)-N-phenylisobutyramide (also known as
22		Isobutyryl Fentanyl).
23	(24)	N-(4-chlorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide (also known
24		as Para-chloroisobutyryl Fentanyl).
25	(25)	N-(4-methoxyphenyl)-N-(1-phenethylpiperidin-4-yl)butyramide (also known
26		as Para-methoxybutyryl Fentanyl).
27	(26)	N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)butyramide (also known as
28		Para-fluorobutyryl Fentanyl).
29	(27)	N-(1-(2-fluorophenethyl)piperidin-4-yl)-N-(2-fluorophenyl)propionamide (also
30		known as 2'-fluoro Ortho-fluorofentanyl; 2'-fluoro 2-fluorofentanyl).

1		(28)	N-(2-methylphenyl)-N-(1-phenethylpiperidin-4-yl)acetamide (also known as
2			Ortho-methyl Acetylfentanyl; 2-methyl acetylfentanyl).
3		(29)	N-(1-phenethylpiperidin-4-yl)-N,3-diphenylpropanamide (also known as
4			Beta'-phenyl Fentanyl <u>: 3-phenylpropanoyl fentanyl</u> and Hydrocinnamoyl
5			Fentanyl).
6		(30)	N-(1-phenethylpiperidin-4-yl)-N-phenylthiophene-2-carboxamide (also
7			known as Thiofuranyl Fentanyl; 2-thiofuranyl fentanyl; thiophene fentanyl).
8		(31)	(E)-N-(1-phenethylpiperidin-4-yl)-N-phenylbut-2-enamide (also known as
9			Crotonyl Fentanyl).
10		<u>(32)</u>	N-(1-(4-methylphenethyl)piperidin-4-yl)-N-phenylacetamide (4'-methyl acetyl
11			fentanyl).
12		(33)	N-phenyl-N-(1-(2-phenylpropyl)piperidin-4-yl)propionamide (beta-methyl
13			fentanyl).
14		(34)	N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)butyramide (ortho-
15			fluorobutyryl fentanyl; 2-fluorobutyryl fentanyl).
16		<u>(35)</u>	2-methoxy-N-(2-methylphenyl)-N-(1-phenethylpiperidin-4-yl)acetamide
17			(ortho-methyl methoxyacetylfentanyl; 2-methyl methoxyacetyl fentanyl).
18		<u>(36)</u>	N-(4-methylphenyl)-N-(1-phenethylpiperidin-4-yl)propionamide (para-
19			methylfentanyl; 4-methylfentanyl).
20		(37)	N-(1-phenethylpiperidin-4-yl)-N-phenylbenzamide (phenyl fentanyl; benzoyl
21			fentanyl).
22		<u>(38)</u>	Ethyl (1-phenethylpiperidin-4-yl)(phenyl)carbamate (fentanyl carbamate).
23		(39)	N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)acrylamide (ortho-fluoroacryl
24			fentanyl).
25		<u>(40)</u>	N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide (ortho-
26			fluoroisobutyryl fentanyl).
27		<u>(41)</u>	N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)furan-2-carboxamide (para-
28			fluoro furanyl fentanyl).
29	4.	Opium d	erivatives. Unless specifically excepted or unless listed in another schedule,
30		any of th	e following opium derivatives, its salts, isomers, and salts of isomers

1 whenever the existence of such salts, isomers, and salts of isomers is possible within 2 the specific chemical designation: 3 a. Acetorphine. 4 b. Acetyldihydrocodeine. 5 Benzylmorphine. C. 6 d. Codeine methylbromide. 7 Codeine-N-Oxide. e. 8 f. Cyprenorphine. 9 Desomorphine. g. 10 h. Dihydromorphine. 11 i. Drotebanol. 12 j. Etorphine (except hydrochloride salt). 13 k. Heroin. 14 I. Hydromorphinol. 15 m. Methyldesorphine. 16 Methyldihydromorphine. n. 17 Morphine methylbromide. 0. 18 p. Morphine methylsulfonate. 19 Morphine-N-Oxide. q. 20 Myrophine. r. 21 S. Nicocodeine. 22 Nicomorphine. t. 23 Normorphine. u. 24 ٧. Pholcodine. 25 W. Thebacon. 26 5. Hallucinogenic substances. Unless specifically excepted or unless listed in another 27 schedule, any material, compound, mixture, or preparation containing any quantity of 28 the following hallucinogenic substances, including their salts, isomers, and salts of 29 isomers whenever the existence of those salts, isomers, and salts of isomers is 30 possible within the specific chemical designation (for purposes of this subsection only, 31 the term "isomer" includes the optical, position, and geometric isomers):

1 Alpha-ethyltryptamine, its optical isomers, salts, and salts of isomers (also known 2 as etryptamine; a-ethyl-1H-indole-3-ethanamine; 3-(2-aminobutyl) indole). 3 b. Alpha-methyltryptamine. 4 4-methoxyamphetamine (also known as 4-methoxy-a-methylphenethylamine; C. 5 paramethoxyamphetamine; PMA). 6 d. N-hydroxy-3,4-methylenedioxyamphetamine (also known as N-hydroxy-alpha-7 methyl-3,4(methylenedioxy)phenylamine, and N-hydroxy MDA. 8 Ibogaine (also known as 7-Ethyl-6, 6B, 7, 8, 9, 10, 12, 13-octahydro-2-methoxye. 9 6, 9-methano-5 H-pyrido [1', 2':1,2] azepino (5,4-b) indole; Tabernanthe iboga). 10 f. Lysergic acid diethylamide. 11 g. Marijuana. 12 Parahexyl (also known as 3-Hexyl-1-hydroxy-7,8,9,10-tetrahydro-6,6,9-trimethylh. 13 6H-dibenzol[b,d]pyran; Synhexyl). 14 Peyote (all parts of the plant presently classified botanically as Lophophora 15 williamsii Lemaire, whether growing or not, the seeds thereof, any extract from 16 any part of such plant, and every compound, manufacture, salts, derivative, 17 mixture, or preparation of such plant, its seeds, or its extracts). 18 j. N-ethyl-3-piperidyl benzilate. 19 k. N-methyl-3-piperidyl benzilate. 20 Psilocybin. Ι. 21 m. (1) Tetrahydrocannabinols, meaning tetrahydrocannabinols naturally contained 22 in a plant of the genus Cannabis (cannabis plant), as well as synthetic 23 equivalents of the substances contained in the cannabis plant, or in the 24 resinous extractives of such plant, including synthetic substances, 25 derivatives, and their isomers with similar chemical structure and 26 pharmacological activity to those substances contained in the plant; such as 27 the following: 28 Delta-1 cis or trans tetrahydrocannabinol, and their optical isomers. (a) 29 Other names: Delta-9-tetrahydrocannabinol. 30 (b) Delta-6 cis or trans tetrahydrocannabinol, and their optical isomers. 31 Other names: Delta-8-tetrahydrocannabinol.

1			(c)	Delta-3,4 cis or trans tetrahydrocannabinol, and its optical isomers.	
2		(Sin	ce no	menclature of these substances is not internationally standardized,	
3		compounds of these structures, regardless of numerical designation of atomic			
4		posi	itions	covered.)	
5		(2)	Tetra	ahydrocannabinols do not include:	
6			(a)	The allowable amount of total tetrahydrocannabinol found in hemp as	
7				defined in chapter 4.1-18.1; or	
8			(b)	A prescription drug approved by the United States food and drug	
9				administration under section 505 of the Federal Food, Drug, and	
10				Cosmetic Act [21 U.S.C. 355].	
11	n.	Can	nabin	oids, synthetic. It includes the chemicals and chemical groups listed	
12		belo	w, inc	cluding their homologues, salts, isomers, and salts of isomers. The term	
13		"iso	mer" i	ncludes the optical, position, and geometric isomers.	
14		(1)	<u>Indo</u>	le acetamides. Any compound structurally derived from 1H-indole3-	
15			<u>acet</u>	amide or 1H-2-acetamide substituted in both of the following ways: at	
16			the r	nitrogen atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl,	
17			<u>alke</u>	nyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2piperidinyl)methyl, 2-	
18			<u>(4-m</u>	orpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-	
19			mor	oholinyl)methyl, tetrahydropyranylmethyl, benzyl, or halo benzyl group;	
20			and,	at the hydrogen of the acetamide by a phenyl, benzyl, cumyl, naphthyl,	
21			<u>adar</u>	mantyl, cyclopropyl, pyrrolidinyl, piperazinyl, or propionaldehyde group	
22			whet	ther or not the compound is further modified to any extent in the	
23			follo	wing ways:	
24			<u>(a)</u>	Substitution to the indole ring to any extent; or	
25			<u>(b)</u>	Substitution to the phenyl, benzyl, cumyl, naphthyl, adamantyl,	
26				cyclopropyl, pyrrolidinyl, piperazinyl, or propionaldehyde group to any	
27				extent; or	
28			<u>(c)</u>	A nitrogen heterocyclic analog of the indole ring; or	
29			<u>(d)</u>	A nitrogen heterocyclic analog of the phenyl, benzyl, naphthyl,	
30				adamantyl, or cyclopropyl ring.	
31			<u>(e)</u>	Examples include:	

1			[1]	N-cyclohexyl-2-(1-pentylindol-3-yl)acetamide - Other names:
2				CH-PIATA, Cyclohexyl-PIATA, CHX-PIATA, CH-PIACA, and
3				CHX-PIACA.
4			<u>[2]</u>	N-cyclohexyl-2-[1-[(4-fluorophenyl)methyl]indol-3-yl]acetamide -
5				Other names: CH-FUBIATA and CH-FUBIACA.
6			<u>[3]</u>	2-[[2-[1-[(4-fluorophenyl)methyl]indol-3-yl]acetyl]amino]-3,3-
7				dimethyl-butanamide - Other names: ADB-FUBIATA, FUB-
8				ACADB, and AD-18.
9	<u>(2)</u>	Indo	le car	boxaldehydes. Any compound structurally derived from 1H-indole-
10		3-ca	rboxa	ldehyde or 1H-2-carboxaldehyde substituted in both of the
11		follo	wing v	ways: at the nitrogen atom of the indole ring by an alkyl, haloalkyl,
12		cyan	oalky	rl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-
13		pipe	ridiny	I)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl,
14		1-(N	-meth	yl-3- morpholinyl)methyl, tetrahydropyranylmethyl, benzyl, or halo
15		benz	yl gro	oup; and, at the hydrogen of the carboxaldehyde by a phenyl,
16		benz	yl, cu	ımyl, naphthyl, adamantyl, cyclopropyl, pyrrolidinyl, piperazinyl, or
17		prop	ional	dehyde group whether or not the compound is further modified to
18		any e	exten	t in the following ways:
19		(a)	Sub	stitution to the indole ring to any extent; or
20		(b)	Sub	stitution to the phenyl, benzyl, cumyl, naphthyl, adamantyl,
21			cycl	opropyl, pyrrolidinyl, piperazinyl, or propionaldehyde group to any
22			exte	ent; or
23		(c)	A ni	trogen heterocyclic analog of the indole ring; or
24		(d)	A ni	trogen heterocyclic analog of the phenyl, benzyl, naphthyl,
25			ada	mantyl, or cyclopropyl ring.
26		(e)	Exa	mples include:
27			[1]	1-Pentyl-3-(1-naphthoyl)indole - Other names: JWH-018 and
28				AM-678.
29			[2]	1-Butyl-3-(1-naphthoyl)indole - Other names: JWH-073.
30			[3]	1-Pentyl-3-(4-methoxy-1-naphthoyl)indole - Other names:
31				JWH-081.

1	[4]	1-[2-(4-morpholinyl)ethyl]-3-(1-naphthoyl)indole - Other names:
2		JWH-200.
3	[5]	1-Propyl-2-methyl-3-(1-naphthoyl)indole - Other names:
4		JWH-015.
5	[6]	1-Hexyl-3-(1-naphthoyl)indole - Other names: JWH-019.
6	[7]	1-Pentyl-3-(4-methyl-1-naphthoyl)indole - Other names:
7		JWH-122.
8	[8]	1-Pentyl-3-(4-ethyl-1-naphthoyl)indole - Other names: JWH-210.
9	[9]	1-Pentyl-3-(4-chloro-1-naphthoyl)indole - Other names:
10		JWH-398.
11	[10]	1-(5-fluoropentyl)-3-(1-naphthoyl)indole - Other names:
12		AM-2201.
13	[11]	1-(2-cyclohexylethyl)-3-(2-methoxyphenylacetyl)indole - Other
14		names: RCS-8.
15	[12]	1-Pentyl-3-(2-methoxyphenylacetyl)indole - Other names:
16		JWH-250.
17	[13]	1-Pentyl-3-(2-methylphenylacetyl)indole - Other names:
18		JWH-251.
19	[14]	1-Pentyl-3-(2-chlorophenylacetyl)indole - Other names: JWH-
20		203.
21	[15]	1-Pentyl-3-(4-methoxybenzoyl)indole - Other names: RCS-4.
22	[16]	(1-(5-fluoropentyl)-3-(2-iodobenzoyl)indole) - Other names:
23		AM-694.
24	[17]	(4-Methoxyphenyl)-[2-methyl-1-(2-(4-morpholinyl)ethyl)indol-3-
25		yl]methanone - Other names: WIN 48,098 and Pravadoline.
26	[18]	(1-Pentylindol-3-yl)-(2,2,3,3-tetramethylcyclopropyl)methanone
27		Other names: UR-144.
28	[19]	(1-(5-fluoropentyl)indol-3-yl)-(2,2,3,3-
29		tetramethylcyclopropyl)methanone - Other names: XLR-11.
30	[20]	(1-(2-morpholin-4-ylethyl)-1H-indol-3-yl)-(2,2,3,3-
31		tetramethylcyclopropyl)methanone - Other names: A-796,260.

1		[[21]	(1-(5-fluoropentyl)-1H-indazol-3-yl)(naphthalen-1-yl)methanone		
2				Other names: THJ-2201.		
3		[[22]	1-naphthalenyl(1-pentyl-1H-indazol-3-yl)-methanone Other		
4				names: THJ-018.		
5		[[23]	(1-(5-fluoropentyl)-1H-benzo[d]imidazol-2-yl)(naphthalen-1-		
6				yl)methanone - Other names: FUBIMINA.		
7		[[24]	1-[(N-methylpiperidin-2-yl)methyl]-3-(adamant-1-oyl) indole -		
8				Other names: AM-1248.		
9		[[25]	1-Pentyl-3-(1-adamantoyl)indole - Other names: AB-001 and		
0				JWH-018 adamantyl analog.		
11	(2) (3)	Indol	le car	boxamides. Any compound structurally derived from 1H-indole-3-		
2		carbo	oxam	ide or 1H-2-carboxamide substituted in both of the following ways:		
3		at the	e nitro	ogen atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl,		
4		alker	nyl, cy	ycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl,		
5		2-(4-	2-(4-morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-			
6		morp	holin	yl)methyl, tetrahydropyranylmethyl, benzyl, or halo benzyl group;		
17		and,	at the	e nitrogen of the carboxamide by a phenyl, benzyl, cumyl,		
8		naph	ithyl,	adamantyl, cyclopropyl, or propionaldehyde group whether or not		
9		the c	ompo	ound is further modified to any extent in the following ways:		
20		(a)	Sub	stitution to the indole ring to any extent; or		
21		(b)	Sub	stitution to the phenyl, benzyl, cumyl, naphthyl, adamantyl,		
22			cycl	opropyl, or propionaldehyde group to any extent; or		
23		(c)	A ni	trogen heterocyclic analog of the indole ring; or		
24		(d)	A ni	trogen heterocyclic analog of the phenyl, benzyl, naphthyl,		
25			ada	mantyl, or cyclopropyl ring.		
26		(e)	Exa	mples include:		
27			[1]	N-Adamantyl-1-pentyl-1H-indole-3-carboxamide - Other names:		
28				JWH-018 adamantyl carboxamide, APICA, SDB-001, and 2NE1.		
29			[2]	N-Adamantyl-1-fluoropentylindole-3-carboxamide - Other names:		
30				STS-135.		

1	[3]	N-Adamantyl-1-pentyl-1H-Indazole-3-carboxamide - Other
2		names: AKB 48 and APINACA.
3	[4]	N-1-naphthalenyl-1-pentyl-1H-indole-3-carboxamide - Other
4		names: NNEI and MN-24.
5	[5]	N-(1-Amino-3,3-dimethyl-1-oxobutan-2-yl)-1-pentyl-1H-indole-3-
6		carboxamide - Other names: ADBICA.
7	[6]	(S)-N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-pentyl-1H-indazole-
8		3-carboxamide - Other names: AB-PINACA.
9	[7]	N-[(1S)-1-(aminocarbonyl)-2-methylpropyl]-1-[(4-
10		fluorophenyl)methyl]-1H-indazole-3-carboxamide - Other names:
11		AB-FUBINACA.
12	[8]	N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(5-fluoropentyl)-1H-
13		indazole-3-carboxamide - Other names: 5-Fluoro AB-PINACA
14		and 5F-AB-PINACA.
15	[9]	N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-pentyl-1H-indazole-
16		3-carboxamide - Other names: ADB-PINACA.
17	[10]	N-[(1S)-1-(aminocarbonyl)-2-methylpropyl]-1-(cyclohexylmethyl)-
18		1H-indazole-3-carboxamide - Other names: AB-CHMINACA.
19	[11]	N-(1-Amino-3,3-dimethyl-1-oxobutan-2-yl)-1-(4-fluorobenzyl)-1H-
20		indazole-3-carboxamide - Other names: ADB-FUBINACA.
21	[12]	N-(adamantan-1-yl)-1-(4-fluorobenzyl)-1H -indazole-3-
22		carboxamide - Other names: FUB-AKB48, FUB-APINACA, and
23		AKB48 N-(4-FLUOROBENZYL).
24	[13]	1-(5-fluoropentyl)-N-(quinolin-8-yl)-1H-indazole-3-carboxamide -
25		Other names: 5-fluoro-THJ.
26	[14]	methyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3-
27		methylbutanoate - Other names: 5-fluoro AMB and 5F-AMB.
28	[15]	methyl 2-(1-(4-fluorobenzyl)-1H-indazole-3-carboxamido)-3-
29		methylbutanoate - Other names: FUB-AMB, MMB-FUBINACA,
30		and AMB-FUBINACA.

1	[16]	N-[1-(aminocarbonyl)-2,2-dimethylpropyl]-1-(cyclohexylmethyl)-1
2		H-indazole-3-carboxamide - Other names: MAB-CHMINACA and
3		ADB-CHMINACA.
4	[17]	Methyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3,3-
5		dimethylbutanoate - Other names: 5F-ADB and
6		5F-MDMB-PINACA.
7	[18]	N-(adamantan-1-yl)-1-(5-fluoropentyl)-1H-indazole-3-
8		carboxamide - Other names: 5F-APINACA and 5F-AKB48.
9	[19]	Methyl 2-(1-(cyclohexylmethyl)-1H-indole-3-carboxamido)-3,3-
10		dimethylbutanoate - Other names: MDMB-CHMICA and
11		MMB-CHMINACA.
12	[20]	Methyl 2-(1-(4-fluorobenzyl)-1H-indazole-3-carboxamido)-3,3-
13		dimethylbutanoate - Other names: MDMB-FUBINACA.
14	[21]	1-(4-cyanobutyl)-N-(2-phenylpropan-2-yl)-1H-indazole-3-carboxa
15		mide - Other names: 4-CN-CUMYL-BUTINACA; 4-cyano-
16		CUMYL-BUTINACA; 4-CN-CUMYL BINACA; CUMYL-4CN
17		-BINACA; SGT-78.
18	[22]	methyl 2-(1-(cyclohexylmethyl)-1H-indole-3-carboxamido)-
19		3-methylbutanoate - Other names: MMB-CHMICA, AMB-
20		CHMICA.
21	[23]	1-(5-fluoropentyl)-N-(2-phenylpropan-2-yl)-1H-pyrrolo[2,3-b]pyridi
22		ne-3-carboxamide - Other names: 5F-CUMYL-P7AICA.
23	[24]	ethyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3,3-
24		dimethylbutanoate - Other names: 5F-EDMB-PINACA.
25	[25]	methyl 2-(1-(5-fluoropentyl)-1H-indole-3-carboxamido)-3,3-
26		dimethylbutanoate - Other names: 5F-MDMB-PICA and 5F-
27		MDMB-2201.
28	[26]	1-(5-fluoropentyl)-N-(2-phenylpropan-2-yl)-1H-indazole-3-
29		carboxamide - Other names: 5F-CUMYL-PINACA, SGT-25.
30	[27]	(1-(4-fluorobenzyl)-1H-indol-3-yl)(2,2,3,3-tetramethylcyclopropyl)
31		methanone - Other names: FUB-144.

1		<u>[28]</u>	methyl 2-(1-(4-fluorobutyl)-1H-indazole-3-carboxamido)-3,3-
2			dimethylbutanoate (4F-MDMB-BINACA, 4F-MDMB-BUTINACA).
3		[29]	Methyl 3,3-dimethyl-2-[(1-pent-4-enylindazole-3-
4			carbonyl)amino]butanoate - Other names: MDMB-4en-PINACA,
5			MDMB-PENINACA, and 5-CL-ADB-A.
6		[30]	Methyl 2-[[1-(5-fluoropentyl)indole-3-carbonyl]amino]-3,3-
7			dimethyl-butanoate - Other names: 5F-MDMB-PICA and 5F-
8			MDMB-2201.
9		<u>[31]</u>	1-butyl-N-(1-carbamoyl-2,2-dimethyl-propyl)indazole-3-
10			carboxamide - Other names: ADB-BINACA and ADB-BUTINACA.
11		[32]	5-bromo-N-(1-carbamoyl-2,2-dimethyl-propyl)-1H-indazole-3-
12			carboxamide - Other names: ADB-5Br-INACA.
13		[33]	Methyl 2-[(5-bromo-1H-indazole-3-carbonyl)amino]-3,3-dimethyl-
14			butanoate - Other names: MDMB-5Br-INACA.
15		[34]	5-bromo-1-butyl-N-(1-carbamoyl-2,2-dimethyl-propyl)indazole-3-
16			carboxamide - Other names: ADB-5'Br-BINACA and ADB-5'Br-
17			BUTINACA.
18	(3) (4)	Indole ca	rboxylic acids. Any compound structurally derived from 1H-indole-
19		3-carbox	ylic acid or 1H-2-carboxylic acid substituted in both of the following
20		ways: at	the nitrogen atom of the indole ring by an alkyl, haloalkyl,
21		cyanoalk	yl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-
22		piperiding	yl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl,
23		1-(N-met	hyl-3- morpholinyl)methyl, tetrahydropyranylmethyl, benzyl, or halo
24		benzyl gr	oup; and, at the hydroxyl group of the carboxylic acid by a phenyl,
25		benzyl, c	umyl, naphthyl, adamantyl, cyclopropyl, or propionaldehyde group
26		whether	or not the compound is further modified to any extent in the
27		following	ways:
28		(a) Su	bstitution to the indole ring to any extent; or
29		(b) Su	bstitution to the phenyl, benzyl, cumyl, naphthyl, adamantyl,
30		сус	clopropyl, propionaldehyde group to any extent; or
31		(c) An	itrogen heterocyclic analog of the indole ring; or

1		(d)	A ni	trogen heterocyclic analog of the phenyl, benzyl, naphthyl,	
2			ada	mantyl, or cyclopropyl ring.	
3		(e)	Exa	mples include:	
4			[1]	1-(cyclohexylmethyl)-1H-indole-3-carboxylic acid 8-quinolinyl	
5				ester - Other names: BB-22 and QUCHIC.	
6			[2]	naphthalen-1-yl 1-(4-fluorobenzyl)-1H-indole-3-carboxylate -	
7				Other names: FDU-PB-22.	
8			[3]	1-pentyl-1H-indole-3-carboxylic acid 8-quinolinyl ester - Other	
9				names: PB-22 and QUPIC.	
10			[4]	1-(5-Fluoropentyl)-1H-indole-3-carboxylic acid 8-quinolinyl ester -	
11				Other names: 5-Fluoro PB-22 and 5F-PB-22.	
12			[5]	quinolin-8-yl-1-(4-fluorobenzyl)-1H-indole-3-carboxylate - Other	
13				names: FUB-PB-22.	
14			[6]	naphthalen-1-yl 1-(5-fluoropentyl)-1H-indole-3-carboxylate -	
15				Other names: NM2201 and CBL2201.	
16	(4) (5)	Napl	hthyln	nethylindoles. Any compound containing a 1H-indol-3-yl-(1-	
17		naph	nthyl)r	methane structure with substitution at the nitrogen atom of the	
18		indo	le ring	g by an alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl,	
19		cyclo	oalkyl	ethyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-	
20		(N-m	nethyl	-2-pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl, or	
21		(tetra	ahydr	opyran-4-yl)methyl group whether or not further substituted in the	
22		indo	le ring	g to any extent and whether or not substituted in the naphthyl ring	
23		to ar	ny ext	ent. Examples include:	
24		(a)	1-P	entyl-1H-indol-3-yl-(1-naphthyl)methane - Other names: JWH-175.	
25		(b)	1-P	entyl-1H-indol-3-yl-(4-methyl-1-naphthyl)methane - Other names:	
26			JWł	- 184.	
27	(5) (6)	Napl	hthoy	lpyrroles. Any compound containing a 3-(1-naphthoyl)pyrrole	
28		struc	cture v	with substitution at the nitrogen atom of the pyrrole ring by an	
29		alkyl	alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-		
30		meth	methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-		
31		pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl, or (tetrahydropyran-4-			

ı		yı)me	etnyl group whether or not further substituted in the pyrrole ring to any
2		exter	nt, whether or not substituted in the naphthyl ring to any extent.
3		Exan	nples include: (5-(2-fluorophenyl)-1-pentylpyrrol-3-yl)-naphthalen-1-
4		ylme	thanone - Other names: JWH-307.
5	(6) (7)	Naph	nthylmethylindenes. Any compound containing a naphthylideneindene
6		struc	ture with substitution at the 3-position of the indene ring by an alkyl,
7		haloa	alkyl, cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-
8		2-pip	eridinyl)methyl, 2 (4 morpholinyl)ethyl, 1-(N-methyl-2-
9		pyrro	olidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl, or (tetrahydropyran-4-
10		yl)me	ethyl group whether or not further substituted in the indene ring to any
11		exter	nt, whether or not substituted in the naphthyl ring to any extent.
12		Exan	nples include: E-1-[1-(1-Naphthalenylmethylene)-1H-inden-3-yl]pentane
13		- Oth	er names: JWH-176.
14	(7) (8)	Cyclo	ohexylphenols. Any compound containing a 2-(3-
15		hydro	oxycyclohexyl)phenol structure with substitution at the 5-position of the
16		phen	olic ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl,
17		cyclo	palkylethyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-
18		(N-m	nethyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl, or
19		(tetra	ahydropyran-4-yl)methyl group whether or not substituted in the
20		cyclo	phexyl ring to any extent. Examples include:
21		(a)	5-(1,1-dimethylheptyl)-2-[(1R,3S)-3-hydroxycyclohexyl]-phenol - Other
22			names: CP 47,497.
23		(b)	5-(1,1-dimethyloctyl)-2-[(1R,3S)-3-hydroxycyclohexyl]-phenol - Other
24			names: Cannabicyclohexanol and CP 47,497 C8 homologue.
25		(c)	5-(1,1-dimethylheptyl)-2-[(1R,2R)-5-hydroxy-2-(3-
26			hydroxypropyl)cyclohexyl]-phenol - Other names: CP 55,940.
27	(8) (9)	Othe	rs specifically named:
28		(a)	(6aR,10aR)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan-2-yl)-
29			6a,7,10,10a-tetrahydrobenzo[c]chromen-1-ol - Other names: HU-210.

1		(b)	(6aS,10aS)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan-2-yl)-	
2			6a,7,10,10a-tetrahydrobenzo[c]chromen-1-ol - Other names:	
3			Dexanabinol and HU-211.	
4		(c)	2,3-Dihydro-5-methyl-3-(4-morpholinylmethyl)pyrrolo[1,2,3-de]-1,4-	
5			benzoxazin-6-yl]-1-napthalenylmethanone - Other names:	
6			WIN 55,212-2.	
7		(d)	Naphthalen-1-yl-(4-pentyloxynaphthalen-1-yl)methanone - Other	
8			names: CB-13.	
9		<u>(e)</u>	N-[(Z)-(1-hexyl-2-oxo-indolin-3-ylidene)amino]benzamide - Other	
10			names: BZO-HEXOXIZID and MDA-19.	
11		<u>(f)</u>	N-[(Z)-(2-oxo-1-pentyl-indolin-3-ylidene)amino]benzamide - Other	
12			names: BZO-POXIZID, Pentyl MDA-19, and 5C-MDA-19.	
13		<u>(g)</u>	N-[(Z)-[1-(5-fluoropentyl)-2-oxo-indolin-3-ylidene]amino]benzamide -	
14			Other names: 5F-BZO-POXIZID and 5F-MDA-19.	
15		<u>(h)</u>	N-[(Z)-(2-oxo-1-pent-4-enyl-indolin-3-ylidene)amino]benzamide -	
16			Other names: BZO-4en-POXIZID and 4en-pentyl MDA-19.	
17		<u>(i)</u>	N-[(Z)-[1-(cyclohexylmethyl)-2-oxo-indolin-3-ylidene]amino]benzamide	
18			- Other names: BZO-CHMOXIZID, Cyclohexylmethyl MDA-19 and	
19			<u>CHM-MDA-19.</u>	
20		<u>(j)</u>	N-(1-carbamoyl-2-methyl-propyl)-2-(5-fluoropentyl)-5-(4-	
21			fluorophenyl)pyrazole-3-carboxamide - Other Names: 5F-AB-	
22			PFUPPYCA.	
23	0.	Substitute	d phenethylamines. This includes any compound, unless specifically	
24		excepted,	specifically named in this schedule, or listed under a different	
25		schedule,	structurally derived from phenylethan-2-amine by substitution on the	
26		phenyl ring in any of the following ways, that is to say, by substitution with a fused		
27		methylene	edioxy ring, fused furan ring, or fused tetrahydrofuran ring; by	
28		substitution with two alkoxy groups; by substitution with one alkoxy and either		
29		one fused furan, tetrahydrofuran, or tetrahydropyran ring system; or by		
30		substitution with two fused ring systems from any combination of the furan,		
31		tetrahydrofuran, or tetrahydropyran ring systems.		

1	(1)	Whether or not the compound is further modified in any of the following	
2		ways	s, that is to say:
3		(a)	By substitution of phenyl ring by any halo, hydroxyl, alkyl,
4			trifluoromethyl, alkoxy, or alkylthio groups;
5		(b)	By substitution at the 2-position by any alkyl groups; or
6		(c)	By substitution at the 2-amino nitrogen atom with alkyl, dialkyl, benzyl,
7			hydroxybenzyl, methylenedioxybenzyl, or methoxybenzyl groups.
8	(2)	Exar	mples include:
9		(a)	2-(4-Chloro-2,5-dimethoxyphenyl)ethanamine (also known as 2C-C or
10			2,5-Dimethoxy-4-chlorophenethylamine).
11		(b)	2-(2,5-Dimethoxy-4-methylphenyl)ethanamine (also known as 2C-D or
12			2,5-Dimethoxy-4-methylphenethylamine).
13		(c)	2-(2,5-Dimethoxy-4-ethylphenyl)ethanamine (also known as 2C-E or
14			2,5-Dimethoxy-4-ethylphenethylamine).
15		(d)	2-(2,5-Dimethoxyphenyl)ethanamine (also known as 2C-H or 2,5-
16			Dimethoxyphenethylamine).
17		(e)	2-(4-lodo-2,5-dimethoxyphenyl)ethanamine (also known as 2C-l or
18			2,5-Dimethoxy-4-iodophenethylamine).
19		(f)	2-(2,5-Dimethoxy-4-nitro-phenyl)ethanamine (also known as 2C-N or
20			2,5-Dimethoxy-4-nitrophenethylamine).
21		(g)	2-(2,5-Dimethoxy-4-(n)-propylphenyl)ethanamine (also known as 2C-
22			P or 2,5-Dimethoxy-4-propylphenethylamine).
23		(h)	2-[4-(Ethylthio)-2,5-dimethoxyphenyl]ethanamine (also known as 2C-
24			T-2 or 2,5-Dimethoxy-4-ethylthiophenethylamine).
25		(i)	2-[4-(Isopropylthio)-2,5-dimethoxyphenyl]ethanamine (also known as
26			2C-T-4 or 2,5-Dimethoxy-4-isopropylthiophenethylamine).
27		(j)	2-(4-bromo-2,5-dimethoxyphenyl)ethanamine (also known as 2C-B or
28			2,5-Dimethoxy-4-bromophenethylamine).
29		(k)	2-(2,5-dimethoxy-4-(methylthio)phenyl)ethanamine (also known as
30			2C-T or 4-methylthio-2,5-dimethoxyphenethylamine).

1	(1)	1-(2,5-dimethoxy-4-iodophenyl)-propan-2-amine (also known as DOI
2		or 2,5-Dimethoxy-4-iodoamphetamine).
3	(m)	1-(4-Bromo-2,5-dimethoxyphenyl)-2-aminopropane (also known as
4		DOB or 2,5-Dimethoxy-4-bromoamphetamine).
5	(n)	1-(4-chloro-2,5-dimethoxy-phenyl)propan-2-amine (also known as
6		DOC or 2,5-Dimethoxy-4-chloroamphetamine).
7	(0)	2-(4-bromo-2,5-dimethoxyphenyl)-N-[(2-
8		methoxyphenyl)methyl]ethanamine (also known as 2C-B-NBOMe;
9		2,5B-NBOMe or 2,5-Dimethoxy-4-bromo-N-(2-
10		methoxybenzyl)phenethylamine).
11	(p)	2-(4-iodo-2,5-dimethoxyphenyl)-N-[(2 -
12		methoxyphenyl)methyl]ethanamine (also known as 2C-I-NBOMe; 2,5I-
13		NBOMe or 2,5-Dimethoxy-4-iodo-N-(2-
14		methoxybenzyl)phenethylamine).
15	(q)	N-(2-Methoxybenzyl)-2-(3,4,5-trimethoxyphenyl)ethanamine (also
16		known as mescaline-NBOMe or 3,4,5-trimethoxy-N-(2-
17		methoxybenzyl)phenethylamine).
18	(r)	2-(4-chloro-2,5-dimethoxyphenyl)-N-[(2-
19		methoxyphenyl)methyl]ethanamine (also known as 2C-C-NBOMe;
20		2,5C-NBOMe or 2,5-Dimethoxy-4-chloro-N-(2-
21		methoxybenzyl)phenethylamine).
22	(s)	2-(7-Bromo-5-methoxy-2,3-dihydro-1-benzofuran-4-yl)ethanamine
23		(also known as 2CB-5-hemiFLY).
24	(t)	2-(8-bromo-2,3,6,7-tetrahydrofuro [2,3-f][1]benzofuran-4-
25		yl)ethanamine (also known as 2C-B-FLY).
26	(u)	2-(10-Bromo-2,3,4,7,8,9-hexahydropyrano[2,3-g]chromen-5-
27		yl)ethanamine (also known as 2C-B-butterFLY).
28	(v)	N-(2-Methoxybenzyl)-1-(8-bromo-2,3,6,7-tetrahydrobenzo[1,2-b:4,5-
29		b']difuran-4-yl)-2-aminoethane (also known as 2C-B-FLY-NBOMe).
30	(w)	1-(4-Bromofuro[2,3-f][1]benzofuran-8-yl)propan-2-amine (also known
31		as bromo-benzodifuranyl-isopropylamine or bromo-dragonFLY).

1		(x)	N-(2-Hydroxybenzyl)-4-iodo-2,5-dimethoxyphenethylamine (also
2			known as 2C-I-NBOH or 2,5I-NBOH).
3		(y)	5-(2-Aminopropyl)benzofuran (also known as 5-APB).
4		(z)	6-(2-Aminopropyl)benzofuran (also known as 6-APB).
5		(aa)	5-(2-Aminopropyl)-2,3-dihydrobenzofuran (also known as 5-APDB).
6		(bb)	6-(2-Aminopropyl)-2,3,-dihydrobenzofuran (also known as 6-APDB).
7		(cc)	2,5-dimethoxy-amphetamine (also known as 2,5-dimethoxy-a-
8			methylphenethylamine; 2,5-DMA).
9		(dd)	2,5-dimethoxy-4-ethylamphetamine (also known as DOET).
10		(ee)	2,5-dimethoxy-4-(n)-propylthiophenethylamine (also known as 2C-T-
11			7).
12		(ff)	5-methoxy-3,4-methylenedioxy-amphetamine.
13		(gg)	4-methyl-2,5-dimethoxy-amphetamine (also known as 4-methyl-2,5-
14			dimethoxy-a-methylphenethylamine; DOM and STP).
15		(hh)	3,4-methylenedioxy amphetamine (also known as MDA).
16		(ii)	3,4-methylenedioxymethamphetamine (also known as MDMA).
17		(jj)	3,4-methylenedioxy-N-ethylamphetamine (also known as N-ethyl-
18			alpha-methyl-3,4(methylenedioxy)phenethylamine, MDE, MDEA).
19		(kk)	3,4,5-trimethoxy amphetamine.
20		(II)	Mescaline (also known as 3,4,5-trimethoxyphenethylamine).
21	p.	Substitute	d tryptamines. This includes any compound, unless specifically
22		excepted,	specifically named in this schedule, or listed under a different
23		schedule,	structurally derived from 2-(1H-indol-3-yl)ethanamine (i.e., tryptamine)
24		by mono-	or di-substitution of the amine nitrogen with alkyl or alkenyl groups or
25		by inclusion	on of the amino nitrogen atom in a cyclic structure whether or not the
26		compound	l is further substituted at the alpha-position with an alkyl group or
27		whether or	not further substituted on the indole ring to any extent with any alkyl,
28		alkoxy, ha	lo, hydroxyl, or acetoxy groups. Examples include:
29		(1) 5-me	thoxy-N,N-diallyltryptamine (also known as 5-MeO-DALT).
30		(2) 4-ace	etoxy-N,N-dimethyltryptamine (also known as 4-AcO-DMT or O-
31		Acety	ylpsilocin).

1 (3) 4-hydroxy-N-methyl-N-ethyltryptamine (also known as 4-HO-MET). 2 4-hydroxy-N,N-diisopropyltryptamine (also known as 4-HO-DIPT). (4) 3 (5)5-methoxy-N-methyl-N-isopropyltryptamine (also known as 5-MeO-MiPT). 4 (6)5-methoxy-N,N-dimethyltryptamine (also known as 5-MeO-DMT). 5 (7) Bufotenine (also known as 3-(Beta-Dimethyl-aminoethyl)-5-hydroxyindole; 6 3-(2-dimethylaminoethyl)-5-indolol; N, N-dimethylserotonin; 5-hydroxy-N,N-7 dimethyltryptamine; mappine). 8 5-methoxy-N,N-diisopropyltryptamine (also known as 5-MeO-DiPT). (8) 9 (9)Diethyltryptamine (also known as N,N-Diethyltryptamine; DET). 10 (10)Dimethyltryptamine (also known as DMT). 11 (11)Psilocyn. 12 1-[3-(trifluoromethylphenyl)]piperazine (also known as TFMPP). q. 13 1-[4-(trifluoromethylphenyl)]piperazine. 14 6,7-dihydro-5H-indeno-(5,6-d)-1,3-dioxol-6-amine (also known as 5,6-S. 15 Methylenedioxy-2-aminoindane or MDAI). 16 2-(Ethylamino)-2-(3-methoxyphenyl)cyclohexanone (also known as 17 Methoxetamine or MXE). 18 u. Ethylamine analog of phencyclidine (also known as N-ethyl-1-19 phenylcyclohexylamine, (1-phenylcyclohexyl) ethylamine, N-(1-phenylcyclohexyl) 20 ethylamine, cyclohexamine, PCE). 21 Pyrrolidine analog of phencyclidine (also known as 1-(1-phenylcyclohexyl)-22 pyrrolidine, PCPy, PHP). 23 Thiophene analog of phencyclidine (also known as (1-[1-(2-thienyl) cyclohexyl] W. 24 piperidine; 2-Thienylanalog of phencyclidine; TPCP, TCP). 25 1-[1-(2-thienyl)cyclohexyl]pyrrolidine (also known as TCPy). Χ. 26 Salvia divinorum, salvinorin A, or any of the active ingredients of salvia divinorum. у. 27 6. Depressants. Unless specifically excepted or unless listed in another schedule, any 28 material compound, mixture, or preparation which contains any quantity of the 29 following substances having a depressant effect on the central nervous system. 30 whenever the existence of such salts, isomers, and salts of isomers is possible within 31 the specific chemical designation:

1 Gamma-hydroxybutyric acid. a. 2 b. Mecloqualone. 3 C. Methaqualone. 4 d. Clonazolam (also known as Clonitrazolam). 5 Etizolam. e. 6 f. Flualprazolam. 7 Flubromazepam. g. 8 Flubromazolam. h. 9 i. Adinazolam. 10 Bromazolam. <u>j.</u> 11 <u>k.</u> <u>Deschloroetizolam.</u> 12 Diclazepam. 13 Stimulants. Unless specifically excepted or unless listed in another schedule, any 14 material, compound, mixture, or preparation which contains any quantity of the 15 following substances having a stimulant effect on the central nervous system. 16 including its salts, isomers, and salts of isomers: 17 Aminorex (also known as 2-amino-5-phenyl-2-oxazoline, or 4,5-dihydro-5-phenyla. 18 2-oxazolamine). 19 Cathinone. b. 20 Substituted cathinones. Any compound, material, mixture, preparation, or other C. 21 product, unless listed in another schedule or an approved food and drug 22 administration drug (e.g., buproprion, pyrovalerone), structurally derived from 2-23 aminopropan-1-one by substitution at the 1-position with either phenyl, naphthyl, 24 or thiophene ring systems, whether or not the compound is further modified in 25 any of the following ways: 26 By substitution in the ring system to any extent with alkyl, alkylenedioxy, (1) 27 alkoxy, haloalkyl, hydroxyl, or halide substituents, whether or not further 28 substituted in the ring system by one or more other univalent substitutents; 29 (2) By substitution at the 3-position with an acyclic alkyl substituent; 30 (3) By substitution at the 2-amino nitrogen atom with alkyl, dialkyl, benzyl, or 31 methoxybenzyl groups; or

1	(4)	By inclusion of the 2-amino nitrogen atom in a cyclic structure.	
2		Some trade or other names:	
3		(a)	3,4-Methylenedioxy-alpha-pyrrolidinopropiophenone (also known as
4			MDPPP).
5		(b)	3,4-Methylenedioxy-N-ethylcathinone (also known as Ethylone,
6			MDEC, or bk-MDEA).
7		(c)	3,4-Methylenedioxy-N-methylcathinone (also known as Methylone or
8			bk-MDMA).
9		(d)	3,4-Methylenedioxypyrovalerone (also known as MDPV).
10		(e)	3,4-Dimethylmethcathinone (also known as 3,4-DMMC).
11		(f)	2-(methylamino)-1-phenylpentan-1-one (also known as Pentedrone).
12		(g)	2-Fluoromethcathinone (also known as 2-FMC).
13		(h)	3-Fluoromethcathinone (also known as 3-FMC).
14		(i)	4-Methylethcathinone (also known as 4-MEC and 4-methyl-N-
15			ethylcathinone).
16		(j)	4-Fluoromethcathinone (also known as Flephedrone and 4-FMC).
17		(k)	4-Methoxy-alpha-pyrrolidinopropiophenone (also known as MOPPP).
18		(I)	4-Methoxymethcathinone (also known as Methedrone; bk-PMMA).
19		(m)	4'-Methyl-alpha-pyrrolidinobutiophenone (also known as MPBP).
20		(n)	Alpha-methylamino-butyrophenone (also known as Buphedrone or
21			MABP).
22		(o)	Alpha-pyrrolidinobutiophenone (also known as alpha-PBP).
23		(p)	Alpha-pyrrolidinopropiophenone (also known as alpha-PPP).
24		(q)	Alpha-pyrrolidinopentiophenone (also known as Alpha-
25			pyrrolidinovalerophenone or alpha-PVP).
26		(r)	Beta-keto-N-methylbenzodioxolylbutanamine (also known as Butylone
27			or bk-MBDB).
28		(s)	Ethcathinone (also known as N-Ethylcathinone).
29		(t)	4-Methylmethcathinone (also known as Mephedrone or 4-MMC).
30		(u)	Methcathinone.
31		(v)	N,N-dimethylcathinone (also known as metamfepramone).

1		(w)	Naphthylpyrovalerone (naphyrone).
2		(x)	B-Keto-Methylbenzodioxolylpentanamine (also known as Pentylone).
3		(y)	4-Methyl-alpha-pyrrolidinopropiophenone (also known as 4-MePPP
4			and MPPP).
5		(z)	1-(1,3-benzodioxol-5-yl)-2-(ethylamino)-pentan-1-one (also known as
6			Ephylone and N-Ethylpentylone).
7		(aa)	N-ethylhexedrone (also known as alpha - ethylaminohexanophenone
8			and 2-(ethylamino)-1-phenylhexan-1-one)).
9		(bb)	Alpha-pyrrolidinohexanophenone (also known as alpha-PHP, alpha-
10			pyrrolidinohexiophenone, and 1-phenyl-2-(pyrrolidin-1-yl)hexan-1-
11			<u>one)</u>).
12		(cc)	4-methyl-alpha-ethylaminopentiophenone (also known as 4-MEAP
13			and 2-(ethylamino)-1-(4-methylphenyl)pentan-1-one)).
14		(dd)	4'-methyl-alpha-pyrrolidinohexiophenone (also known as MPHP <u>, 4'-</u>
15			methyl-alpha-pyrrolidinohexanophenone and 1-(4-methylphenyl)-2-
16			(pyrrolidin-1-yl)hexan-1-one)).
17		(ee)	Alpha-pyrrolidinoheptaphenone (also known as PV8 and 1-phenyl-2-
18			(pyrrolidin-1-yl)heptan-1-one)).
19		(ff)	4-chloro-alpha-pyrrolidinovalerophenone (also known 4-chloro-alpha-
20			PVP, 4'-chloro-alpha-pyrrolidinopentiophenone, and 1-(4-
21			chlorophenyl)-2-(pyrrolidin-1-yl)pentan-1-one)).
22	d.	Fenethylli	ne.
23	e.	Fluoroam	phetamine.
24	f.	Fluorome	thamphetamine.
25	g.	(±)cis-4-m	nethylaminorex (also known as (±)cis-4,5-dihydro-4-methyl-5-phenyl-2-
26		oxazolam	ine).
27	h.	N-Benzylp	piperazine (also known as BZP, 1-benzylpiperazine).
28	i.	N-ethylamphetamine.	
29	j.	N, N-dime	ethylamphetamine (also known as N,N-alpha-trimethyl-
30		benzenee	thanamine; N,N-alpha-trimethylphenethylamine).

1		k.	1-(4-methoxyphenyl)-N-methylpropan-2-amine (also known as			
2			paramethoxymethamphetamine and PMMA).			
3		<u>l.</u>	4,4'-Dimethylaminorex (4,4'-DMAR; 4,5-dihydro-4-methyl-5-(4-methylphenyl)-2-			
4			oxazolamine; 4-methyl-5-(4-methylphenyl)-4,5-dihydro-1,3-oxazol-2-amine).			
5		<u>m.</u>	Amineptine (Also known as 7- [(10,11-dihydro-5Hdibenzo[a,d]cyclohepten-5-			
6			yl)amino]heptanoic acid).			
7		<u>n.</u>	Mesocarb (Also known as N-phenyl-N' -(3-(1- phenylpropan-2-yl)-1,2,3-			
8			oxadiazol-3- ium-5-yl)carbamimidate).			
9		<u>o.</u>	Methiopropamine (Also known as N-methyl-1-(thiophen-2-yl)propan-2-amine).			
10	SEC	OIT	2. AMENDMENT. Section 19-03.1-11 of the North Dakota Century Code is			
11	amende	d and	d reenacted as follows:			
12	19-0	3.1-1	11. Schedule IV.			
13	1.	The	controlled substances listed in this section are included in schedule IV.			
14	2.	Sch	edule IV consists of the drugs and other substances, by whatever official name,			
15		com	common or usual name, chemical name, or brand name designated, listed in this			
16		sec	section.			
17	3.	Nar	cotic drugs. Unless specifically excepted or unless listed in another schedule, any			
18		mat	erial, compound, mixture, or preparation containing any of the following narcotic			
19		drug	gs or their salts calculated as the free anhydrous base or alkaloid, in limited			
20		qua	ntities as set forth below:			
21		a.	Not more than 1 milligram of difenoxin and not less than 25 micrograms of			
22			atropine sulfate per dosage unit.			
23		b.	Dextropropoxyphene (also known as alpha-(+)-4-dimethylamino- 1,2-diphenyl-3-			
24			methyl-2-propionoxybutane).			
25		C.	2-[(dimethylamino)methyl]-1-(3-methoxyphenyl)cyclohexanol, its salts, optical			
26			and geometric isomers and salts of these isomers including Tramadol.			
27	4.	Dep	ressants. Unless specifically excepted or unless listed in another schedule, any			
28		mat	erial, compound, mixture, or preparation containing any quantity of the following			
29		sub	stances, including their salts, isomers, and salts of isomers whenever the			
30		exis	tence of those salts, isomers, and salts of isomers is possible within the specific			
31		che	mical designation:			

1	a.	Alprazolam.
2	b.	Alfaxalone.
3	C.	Barbital.
4	d.	Brexanolone.
5	e.	Bromazepam.
6	f.	Camazepam.
7	g.	Carisoprodol.
8	h.	Chloral betaine.
9	i.	Chloral hydrate.
10	j.	Chlordiazepoxide.
11	k.	Clobazam.
12	l.	Clonazepam.
13	m.	Clorazepate.
14	n.	Clotiazepam.
15	0.	Cloxazolam.
16	<u>p.</u>	Daridorexant.
17	p. <u>q.</u>	Delorazepam.
18	q. <u>r.</u>	Diazepam.
19	r. s.	Dichloralphenazone.
20	s. <u>t.</u>	Estazolam.
21	t. <u>u.</u>	Ethchlorvynol.
22	u. v.	Ethinamate.
23	∀. <u>W.</u>	Ethyl loflazepate.
24	₩. <u>X.</u>	Fludiazepam.
25	Х. <u>У.</u>	Flunitrazepam.
26	y. z.	Flurazepam.
27	z. aa.	Fospropofol.
28	aa. bb.	Halazepam.
29	bb. cc.	Haloxazolam.
30	cc. dd.	Indiplon.
31	dd. ee.	Ketazolam.

1 ee.ff. Lemborexant. 2 ff.gg. Loprazolam. 3 gg.hh. Lorazepam. 4 hh.ii. Lorcaserin. 5 ||.jj. Lormetazepam. 6 jj.kk. Mebutamate. 7 kk.II. Medazepam. 8 Meprobamate. ll.mm. 9 mm.nn. Methohexital. 10 Methylphenobarbital (also known as mephobarbital). nn.00. 11 Midazolam. 00.pp. 12 pp.qq. Nimetazepam. 13 Nitrazepam. qq.rr. 14 Nordiazepam. ff.ss. 15 ss.tt. Oxazepam. 16 Oxazolam. tt.<u>uu.</u> 17 uu.vv. Paraldehyde. 18 ww.ww. Petrichloral. 19 ww.xx. Phenobarbital. 20 Pinazepam. ХХ.<u>УУ.</u> 21 yy.<u>zz.</u> Propofol. 22 zz.aaa. Prazepam. 23 aaa.bbb. Quazepam. 24 bbb.ccc. Remimazolam. 25 Suvorexant. ccc.ddd. 26 ddd.eee. Temazepam. 27 eee.fff. Tetrazepam. 28 fff.ggg. Triazolam. 29 Zaleplon. ggg.hhh. 30 hhh.iii. Zolpidem. 31 |||.|||<u>|</u>. Zopiclone.

1 Fenfluramine. Any material, compound, mixture, or preparation which contains any 2 quantity of the following substances, including its salts, isomers (whether optical, 3 position, or geometric), and salts of such isomers, whenever the existence of such 4 salts, isomers, and salts of isomers is possible: Fenfluramine. 5 6.5. Stimulants. Unless specifically excepted or unless listed in another schedule, any 6 material, compound, mixture, or preparation which contains any quantity of the 7 following substances having a stimulant effect on the central nervous system, 8 including its salts, isomers, and salts of isomers: 9 a. Cathine. 10 b. Diethylpropion. 11 C. Fencamfamin. 12 d. Fenproporex. 13 Mazindol. e. 14 f. Mefenorex. 15 g. Modafinil. 16 Pemoline (including organometallic complexes and chelates thereof). h. 17 i. Phentermine. 18 j. Pipradrol. 19 k. Serdexmethylphenidate. 20 Sibutramine. <u>l.</u> 21 l.m. Solriamfetol. 22 SPA ((-)-1-dimethylamino-1, 2-diphenylethane). m.n. 23 Other substances. Unless specifically excepted or unless listed in another schedule, 7.6. 24 any material, compound, mixture, or preparation which contains any quantity of: 25 a. Pentazocine, including its salts. 26 Butorphanol, including its optical isomers. b. 27 Eluxadoline (5-[[(2S)-2-amino-3-[4-aminocarbonyl)-2,6-dimethylphenyl]-1-C. 28 oxopropyl][(1S)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]amino]methyl]-2-29 methoxybenzoic acid) (including its optical isomers) and its salts, isomers, and 30 salts of isomers.

- 1 The board may except by rule any compound, mixture, or preparation containing any 8.7. 2 depressant substance listed in subsection 2 from the application of all or any part of 3 this chapter if the compound, mixture, or preparation contains one or more active 4 medicinal ingredients not having a depressant effect on the central nervous system, 5 and if the admixtures are included therein in combinations, quantity, proportion, or 6 concentration that vitiate the potential for abuse of the substances which have a 7 depressant effect on the central nervous system.
- 8 **SECTION 3. AMENDMENT.** Section 19-03.1-13 of the North Dakota Century Code is 9 amended and reenacted as follows:

19-03.1-13. Schedule V.

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- 1. The controlled substances listed in this section are included in schedule V.
- 12 2. Schedule V consists of the drugs and other substances, by whatever official name, 13 common or usual name, chemical name, or brand name designated, listed in this section.
 - 3. Narcotic drugs. Unless specifically excepted or unless listed in another schedule, any material, compound, mixture, or preparation containing any of the following narcotic drugs and their salts.
 - Narcotic drugs containing non-narcotic active medicinal ingredients. Any compound, mixture, or preparation containing any of the following narcotic drugs, or their salts calculated as the free anhydrous base or alkaloid, in limited quantities as set forth below, which includes one or more non-narcotic active medicinal ingredients in sufficient proportion to confer upon the compound, mixture, or preparation valuable medicinal qualities other than those possessed by narcotic drugs alone.
 - a. Not more than 200 milligrams of codeine per 100 milliliters or per 100 grams.
 - Not more than 100 milligrams of dihydrocodeine per 100 milliliters or per b. 100 grams.
 - Not more than 100 milligrams of ethylmorphine per 100 milliliters or per 100 grams.
 - Ganaxolone (3alpha-hydroxy-3beta-methyl-5alpha-pregnan-20-one). d.
 - Not more than 2.5 milligrams of diphenoxylate and not less than 25 micrograms <u>e.</u> of atropine sulfate per dosage unit.

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1 e.f. Not more than 100 milligrams of opium per 100 milliliters or per 100 grams. 2 Not more than 0.5 milligram of difenoxin and not less than 25 micrograms of f.g. 3 atropine sulfate per dosage unit. 4 5. Depressants. Unless specifically exempted or excluded or unless listed in another 5 schedule, any material, compound, mixture, or preparation that contains any quantity 6 of the following substances having a depressant effect on the central nervous system. 7 including its salts, isomers, and salts of isomers whenever the existence of such salts, 8 isomers, and salts of isomers is possible: 9 Brivaracetam ((2S)-2-[(4R)-2-oxo-4-propylpyrrolidin-1-yl]butanamide) (also a. 10 referred to as BRV; UCB-34714; Briviact) (including its salts). 11 Cenobamate [(1R)-1-(2-chlorophenyl)-2-(tetrazol-2-yl)ethyl] carbamate; 2Hb. 12 tetrazole-2-ethanol, alpha-(2-chlorophenyl)-, carbamate (ester), (alphaR)-; 13 carbamic acid (R)-(+)-1-(2-chlorophenyl)-2-(2H-tetrazol-2-yl)ethyl ester). 14 Ezogabine N-[2-amino-4-(4-fluorobenzylamino)-phenyl]-carbamic acid ethyl ester. C. 15 d. Lacosamide [(R)-2-acetoamido-N-benzyl-3-methoxy-propionamide]. 16 Lasmiditan [2,4,6-trifluoro-N-(6-(1-methylpiperidine-4-carbonyl)pyridine-2-yle. 17 benzamide]. 18 f. Pregabalin [(S)-3-(aminomethyl)-5-methylhexanoic acid]. 19 Gabapentin [2-[1-(aminomethyl) cyclohexyl] acetic acid]. g. 20 6. Stimulants. Unless specifically exempted or excluded or unless listed in another 21 schedule, any material, compound, mixture, or preparation containing any quantity of 22 the following substances having a stimulant effect on the central nervous system, 23 including their salts, isomers, and salts of isomers: Pyrovalerone.

SECTION 4. EMERGENCY. This Act is declared to be an emergency measure.