Sixty-sixth Legislative Assembly of North Dakota

HOUSE BILL NO. 1113

Introduced by

Judiciary Committee

(At the request of the State Board of Pharmacy)

- 1 A BILL for an Act to amend and reenact subsection 18 of section 19-03.1-01, section
- 2 19-03.1-05, subsection 7 of section 19-03.1-07, subsection 4 of section 19-03.1-09,
- 3 subsection 7 of section 19-03.1-11, and subsection 5 of section 19-03.1-13 of the North Dakota
- 4 Century Code, relating to the definition of marijuana and the scheduling of controlled
- 5 substances; and to declare an emergency.

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

7 SECTION 1. AMENDMENT. Subsection 18 of section 19-03.1-01 of the North Dakota

8 Century Code is amended and reenacted as follows:

- 9 18. "Marijuana" means all parts of the plant cannabis <u>sativa L.</u>, whether growing or not;
- 10 the seeds thereof; the resinous product of the combustionresin extracted from any part
- 11 of the plant cannabis; and every compound, manufacture, salt, derivative, mixture, or

12 preparation of the plant or, its seeds, <u>or resin</u>. The term does not include the mature

- 13 stalks of the plant, fiber produced from the stalks, oil or cake made from the seeds of
- 14 the plant, any other compound, manufacture, salt, derivative, mixture, or preparation of
- mature stalks, <u>except the resin extracted therefrom</u>, fiber, oil, or cake, or the sterilized
 seed of the plant which is incapable of germination.
- SECTION 2. AMENDMENT. Section 19-03.1-05 of the North Dakota Century Code is
 amended and reenacted as follows:
- 19 **19-03.1-05. Schedule I.**
- 20 1. The controlled substances listed in this section are included in schedule I.
- Schedule I consists of the drugs and other substances, by whatever official name,
 common or usual name, chemical name, or brand name designated, listed in this
 section.

1	3.	Opi	ates. Unless specifically excepted or unless listed in another schedule, any of the					
2		follo	following opiates, including their isomers, esters, ethers, salts, and salts of isomers,					
3		este	esters, and ethers, whenever the existence of those isomers, esters, ethers, and salts					
4		is p	ossible within the specific chemical designation:					
5		a.	Acetylmethadol.					
6		b.	Allylprodine.					
7		C.	Alphacetylmethadol.					
8		d.	Alphameprodine.					
9		e.	Alphamethadol.					
10		f.	Benzethidine.					
11		g.	Betacetylmethadol.					
12		h.	Betameprodine.					
13		i.	Betamethadol.					
14		j.	Betaprodine.					
15		k.	Clonitazene.					
16		I.	Dextromoramide.					
17		m.	Diampromide.					
18		n.	Diethylthiambutene.					
19		0.	Difenoxin.					
20		p.	Dimenoxadol.					
21		q.	Dimepheptanol.					
22		r.	Dimethylthiambutene.					
23		S.	Dioxaphetyl butyrate.					
24		t.	Dipipanone.					
25		u.	Ethylmethylthiambutene.					
26		V.	Etonitazene.					
27		W.	Etoxeridine.					
28		Х.	Furethidine.					
29		у.	Hydroxypethidine.					
30		Z.	Ketobemidone.					
31		aa.	Levomoramide.					

1	bb.	Levophenacylmorphan.
2	CC.	Morpheridine.
3	dd.	MPPP (also known as 1-methyl-4-phenyl-4-propionoxypiperidine).
4	ee.	Noracymethadol.
5	ff.	Norlevorphanol.
6	gg.	Normethadone.
7	hh.	Norpipanone.
8	ii.	PEPAP (1-(2-Phenylethyl)-4-Phenyl-4-acetoxypiperidine).
9	jj.	Phenadoxone.
10	kk.	Phenampromide.
11	Ш.	Phenomorphan.
12	mm.	Phenoperidine.
13	nn.	Piritramide.
14	00.	Proheptazine.
15	pp.	Properidine.
16	qq.	Propiram.
17	rr.	Racemoramide.
18	SS.	Tilidine.
19	tt.	Trimeperidine.
20	uu.	3,4-dichloro-N-[2-(dimethylamino)cyclbhexylcyclohexyl]-N-methylbenzamide
21		(also known as U-47700).
22	VV.	1-cyclohexyl-4-(1,2-diphenylethyl)piperazine (also know as MT-45).
23	WW.	3,4-dichloro-N-{[1-(dimethylamino)cyclohexyl]methyl}benzamide (also known as
24		AH-7921).
25	XX.	Fentanyl derivatives. Unless specifically excepted or unless listed in another
26		schedule or are not FDA approved drugs, and are derived from N-(1-(2-
27		Phenylethyl)-4-piperidinyl)-N-phenylpropanamide (Fentanyl) by any substitution
28		on or replacement of the phenethyl group, any substitution on the piperidine ring,
29		any substitution on or replacement of the propanamide group, any substitution on
30		the anilido phenyl group, or any combination of the above. Examples include:

1	(1)	N-[1-(1-methyl-2-phenethyl)-4-piperidinyl]-N-phenylacetamide (also known
2		as Acetyl-alpha-methylfentanyl).
3	(2)	N-[1-(alpha-methyl-beta-phenyl)ethyl-4-piperidyl]propionanilide; 1-(1-methyl-
4		2-phenylethyl)-4-(N-propanilido)piperidine (also known as Alpha-
5		methylfentanyl).
6	(3)	N-[1-methyl-2-(2-thienyl)ethyl-4-piperidinyl]-N-phenylpropanamide (also
7		known as Alpha-methylthiofentanyl).
8	(4)	N-[1-(2-hydroxy-2-phenethyl)-4-piperidinyl]-N-phenylpropanamide (also
9		known as Beta-hydroxyfentanyl).
10	(5)	N-[1-(2-hydroxy-2-phenethyl)-3-methyl-4-piperidinyl]-N-phenylpropanamide
11		(also known as Beta-hydroxy-3-methylfentanyl).
12	(6)	N-[3-methyl-1-(2-phenylethyl)-4-piperidyl]-N-phenylpropanamide (also
13		known as 3-Methylfentanyl).
14	(7)	N-[3-methyl-1-(2-thienyl)ethyl-4-piperidinyl]-N-phenylpropanamide (also
15		known as 3-Methylthiofentanyl).
16	(8)	N-(4-fluorophenyl)-N-[1-(2-phenethyl)-4-piperidinyl]propanamide (also
17		known as Para-fluorofentanyl).
18	(9)	N-phenyl-N-[1-(2-thienyl)ethyl-4-piperidinyl]propanamide (also known as
19		Thiofentanyl).
20	(10)	N-(1-phenylethylpiperidin-4-yl)-N-phenylfuran-2-carboxamide (also known
21		as Furanyl Fentanyl).
22	(11)	N-(1-phenethylpiperidin-4-yl)-N-phenylbutyramide; N-(1-phenethylpiperidin-
23		4-yl)-N-phenylbutanamide (also known as Butyryl Fentanyl).
24	(12)	N-[1-[2-hydroxy-2-(thiophen-2-yl)ethyl]piperidin-4-yl]-N-phenylpropionamide;
25		N-[1-[2-hydroxy-2-(2-thienyl)ethyl]-4-piperidinyl]-N-phenylpropanamide (also
26		known as Beta-Hydroxythiofentanyl).
27	(13)	N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide (also known as Acetyl
28		Fentanyl).
29	(14)	N-phenyl-N-[1-(2-phenylethyl)piperidin-4-yl]prop-2-enamideN-(1-phenethylpi
30		peridin-4-yl)-N-phenylacrylamide (also known as AcrylfentanylAcryl
31		<u>Fentanyl</u>).

1		(15)	N-phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]-pentanamideN-(1-phenethylpip
2			<u>eridin-4-yl)-N-phenylpentanamide</u> (also known as Valeryl Fentanyl).
3		<u>(16)</u>	N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide (also known
4			as 4-Fluoroisobutyryl Fentanyl).
5		<u>(17)</u>	N-(2-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)propionamide (also known
6			as Ortho-fluorofentanyl, 2-Fluorofentanyl).
7		<u>(18)</u>	N-(1-phenethylpiperidin-4-yl)-N-phenyltetrahydrofuran-2-carboxamide (also
8			known as Tetrahydrofuranyl Fentanyl).
9		<u>(19)</u>	2-methoxy-N-(1-phenethylpiperidin-4-yl)-N-phenylacetamide (also known as
10			Methoxyacetyl Fentanyl).
11		<u>(20)</u>	N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopropanecarboxamide (also
12			known as Cyclopropyl Fentanyl).
13		<u>(21)</u>	N-(2-fluorophenyl)-2-methoxy-N-(1-phenethylpiperidin-4-yl)acetamide (also
14			known as Ocfentanil).
15		<u>(22)</u>	N-(1-phenethylpiperidin-4-yl)-N-phenylcyclopentanecarboxamide (also
16			known as Cyclopentyl Fentanyl).
17		<u>(23)</u>	N-(1-phenethylpiperidin-4-yl)-N-phenylisobutyramide (also known as
18			Isobutyryl Fentanyl).
19		<u>(24)</u>	N-(4-chlorophenyl)-N-(1-phenethylpiperidin-4-yl)isobutyramide (also known
20			as Para-chloroisobutyryl Fentanyl).
21		<u>(25)</u>	N-(4-methoxyphenyl)-N-(1-phenethylpiperidin-4-yl)butyramide (also known
22			as Para-methoxybutyryl Fentanyl).
23		<u>(26)</u>	N-(4-fluorophenyl)-N-(1-phenethylpiperidin-4-yl)butyramide (also known as
24			Para-fluorobutyryl Fentanyl).
25	4.	Opium d	erivatives. Unless specifically excepted or unless listed in another schedule,
26		any of th	e following opium derivatives, its salts, isomers, and salts of isomers
27		wheneve	er the existence of such salts, isomers, and salts of isomers is possible within
28		the spec	ific chemical designation:
29		a. Ace	etorphine.
30		b. Ace	etyldihydrocodeine.
31		c. Ber	zylmorphine.

1		d.	Codeine methylbromide.				
2		e.	Codeine-N-Oxide.				
3		f.	Cyprenorphine.				
4		g.	Desomorphine.				
5		h.	Dihydromorphine.				
6		i.	Drotebanol.				
7		j.	Etorphine (except hydrochloride salt).				
8		k.	Heroin.				
9		I.	Hydromorphinol.				
10		m.	Methyldesorphine.				
11		n.	Methyldihydromorphine.				
12		0.	Morphine methylbromide.				
13		p.	Morphine methylsulfonate.				
14		q.	Morphine-N-Oxide.				
15		r.	Myrophine.				
16		S.	Nicocodeine.				
17		t.	Nicomorphine.				
18		u.	Normorphine.				
19		V.	Pholcodine.				
20		W.	Thebacon.				
21	5.	Hal	lucinogenic substances. Unless specifically excepted or unless listed in another				
22		sch	edule, any material, compound, mixture, or preparation containing any quantity of				
23		the	following hallucinogenic substances, including their salts, isomers, and salts of				
24		isor	mers whenever the existence of those salts, isomers, and salts of isomers is				
25		pos	sible within the specific chemical designation (for purposes of this subsection only,				
26		the	term "isomer" includes the optical, position, and geometric isomers):				
27		a.	Alpha-ethyltryptamine, its optical isomers, salts, and salts of isomers (also known				
28			as etryptamine; a-ethyl-1H-indole-3-ethanamine; 3-(2-aminobutyl) indole).				
29		b.	Alpha-methyltryptamine.				
30		C.	4-methoxyamphetamine (also known as 4-methoxy-a-methylphenethylamine;				
31			paramethoxyamphetamine; PMA).				

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

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1		0.	Cann	abinoi	ls, synthetic. It includes the chemicals and chemical groups listed
2			below	v, inclu	ling their homologues, salts, isomers, and salts of isomers. The term
3			"isom	er" inc	udes the optical, position, and geometric isomers.
4			(1)	Indole	carboxaldehydes. Any compound structurally derived from 1H-indole-
5			:	3-carb	oxaldehyde or 1H-2-carboxaldehyde substituted in both of the
6			1	followi	ng ways: at the nitrogen atom of the indole ring by an alkyl, haloalkyl,
7			(cyanoa	lkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-
8			l	piperid	nyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl,
9				1-(N-m	ethyl-3- morpholinyl)methyl, tetrahydropyranylmethyl, benzyl, or halo
10			I	benzyl	group; and, at the hydrogen of the carboxaldehyde by a phenyl,
11			I	benzyl	cumyl, naphthyl, adamantyl, cyclopropyl, pyrrolidinyl, piperazinyl, or
12			I	propio	aldehyde group whether or not the compound is further modified to
13			i	any ex	ent in the following ways:
14				(a) S	substitution to the indole ring to any extent; or
15				(b) S	substitution to the phenyl, benzyl, <u>cumyl,</u> naphthyl, adamantyl,
16				C	yclopropyl, <u>pyrrolidinyl, piperazinyl,</u> or propionaldehyde group to any
17				e	xtent; or
18				(c) A	nitrogen heterocyclic analog of the indole ring; or
19				(d) A	nitrogen heterocyclic analog of the phenyl, benzyl, naphthyl,
20				a	damantyl, or cyclopropyl ring.
21				(e) E	xamples include:
22				[′] 1-Pentyl-3-(1-naphthoyl)indole - Other names: JWH-018 and
23					AM-678.
24				[2] 1-Butyl-3-(1-naphthoyl)indole - Other names: JWH-073.
25				[:] 1-Pentyl-3-(4-methoxy-1-naphthoyl)indole - Other names:
26					JWH-081.
27				[4] 1-[2-(4-morpholinyl)ethyl]-3-(1-naphthoyl)indole - Other names:
28					JWH-200.
29				[<u></u> {] 1-Propyl-2-methyl-3-(1-naphthoyl)indole - Other names:
30					JWH-015.
31				[6] 1-Hexyl-3-(1-naphthoyl)indole - Other names: JWH-019.

1	[7]	1-Pentyl-3-(4-methyl-1-naphthoyl)indole - Other names:
2	[1]	JWH-122.
3	[8]	1-Pentyl-3-(4-ethyl-1-naphthoyl)indole - Other names: JWH-210.
4	[9]	1-Pentyl-3-(4-chloro-1-naphthoyl)indole - Other names:
5		JWH-398.
6	[10]	1-(5-fluoropentyl)-3-(1-naphthoyl)indole - Other names:
7		AM-2201.
8	[11]	1-(2-cyclohexylethyl)-3-(2-methoxyphenylacetyl)indole - Other
9		names: RCS-8.
10	[12]	1-Pentyl-3-(2-methoxyphenylacetyl)indole - Other names:
11		JWH-250.
12	[13]	1-Pentyl-3-(2-methylphenylacetyl)indole - Other names:
13		JWH-251.
14	[14]	1-Pentyl-3-(2-chlorophenylacetyl)indole - Other names: JWH-
15		203.
16	[15]	1-Pentyl-3-(4-methoxybenzoyl)indole - Other names: RCS-4.
17	[16]	(1-(5-fluoropentyl)-3-(2-iodobenzoyl)indole) - Other names:
18		AM-694.
19	[17]	(4-Methoxyphenyl)-[2-methyl-1-(2-(4-morpholinyl)ethyl)indol-3-
20		yl]methanone - Other names: WIN 48,098 and Pravadoline.
21	[18]	(1-Pentylindol-3-yl)-(2,2,3,3-tetramethylcyclopropyl)methanone
22		Other names: UR-144.
23	[19]	(1-(5-fluoropentyl)indol-3-yl)-(2,2,3,3-
24		tetramethylcyclopropyl)methanone - Other names: XLR-11.
25	[20]	(1-(2-morpholin-4-ylethyl)-1H-indol-3-yl)-(2,2,3,3-
26		tetramethylcyclopropyl)methanone - Other names: A-796,260.
27	[21]	(1-(5-fluoropentyl)-1H-indazol-3-yl)(naphthalen-1-yl)methanone
28		Other names: THJ-2201.
29	[22]	1-naphthalenyl(1-pentyl-1H-indazol-3-yl)-methanone Other
30		names: THJ-018.

1		[[23]	(1-(5-fluoropentyl)-1H-benzo[d]imidazol-2-yl)(naphthalen-1-
2				yl)methanone - Other names: FUBIMINA.
3		[[24]	1-[(N-methylpiperidin-2-yl)methyl]-3-(adamant-1-oyl) indole -
4				Other names: AM-1248.
5		[25]	1-Pentyl-3-(1-adamantoyl)indole - Other names: AB-001 and
6				JWH-018 adamantyl analog.
7	(2)	Indol	le car	boxamides. Any compound structurally derived from 1H-indole-3-
8		carbo	oxam	ide or 1H-2-carboxamide substituted in both of the following ways:
9		at the	e nitro	ogen atom of the indole ring by an alkyl, haloalkyl, cyanoalkyl,
10		alker	nyl, c	vcloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl,
11		2-(4-	morp	holinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-
12		morp	holin	yl)methyl, tetrahydropyranylmethyl, benzyl, or halo benzyl group;
13		and,	at the	e nitrogen of the carboxamide by a phenyl, benzyl, <u>cumyl,</u>
14		naph	ithyl,	adamantyl, cyclopropyl, or propionaldehyde group whether or not
15		the c	ompo	ound is further modified to any extent in the following ways:
16		(a)	Sub	stitution to the indole ring to any extent; or
17		(b)	Sub	stitution to the phenyl, benzyl, <u>cumyl,</u> naphthyl, adamantyl,
18			cycl	opropyl, or propionaldehyde group to any extent; or
19		(C)	A ni	trogen heterocyclic analog of the indole ring; or
20		(d)	A ni	trogen heterocyclic analog of the phenyl, benzyl, naphthyl,
21			ada	mantyl, or cyclopropyl ring.
22		(e)	Exa	mples include:
23			[1]	N-Adamantyl-1-pentyl-1H-indole-3-carboxamide - Other names:
24				JWH-018 adamantyl carboxamide, APICA, SDB-001, and 2NE1.
25			[2]	N-Adamantyl-1-fluoropentylindole-3-carboxamide - Other names:
26				STS-135.
27			[3]	N-Adamantyl-1-pentyl-1H-Indazole-3-carboxamide - Other
28				names: AKB 48 and APINACA.
29			[4]	N-1-naphthalenyl-1-pentyl-1H-indole-3-carboxamide - Other
30				names: NNEI and MN-24.

1	[5]	N-(1-Amino-3,3-dimethyl-1-oxobutan-2-yl)-1-pentyl-1H-indole-3-
2		carboxamide - Other names: ADBICA.
3	[6]	(S)-N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-pentyl-1H-indazole-
4		3-carboxamide - Other names: AB-PINACA.
5	[7]	N-[(1S)-1-(aminocarbonyl)-2-methylpropyl]-1-[(4-
6		fluorophenyl)methyl]-1H-indazole-3-carboxamide - Other names:
7		AB-FUBINACA.
8	[8]	(S)-N-(1-amino-3-methyl-1-oxobutan-2-yl)-1-(5-fluoropentyl)-1H-
9		indazole-3-carboxamide - Other names: 5-Fluoro AB-PINACA
10		and 5F-AB-PINACA.
11	[9]	N-(1-amino-3,3-dimethyl-1-oxobutan-2-yl)-1-pentyl-1H-indazole-
12		3-carboxamide - Other names: ADB-PINACA.
13	[10]	N-[(1S)-1-(aminocarbonyl)-2-methylpropyl]-1-(cyclohexylmethyl)-
14		1H-indazole-3-carboxamide - Other names: AB-CHMINACA.
15	[11]	N-(1-Amino-3,3-dimethyl-1-oxobutan-2-yl)-1-(4-fluorobenzyl)-1H-
16		indazole-3-carboxamide - Other names: ADB-FUBINACA.
17	[12]	N-((3s,5s,7s)-adamantan-1-yl)-1-(4-fluorobenzyl)-1H-indazole-3-
18		carboxamide - Other names: FUB-AKB48 and AKB48 N-(4-
19		fluorobenzyl) analog.
20	[13]	1-(5-fluoropentyl)-N-(quinolin-8-yl)-1H-indazole-3-carboxamide -
21		Other names: 5-fluoro-THJ.
22	[14]	(S)-methyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3-
23		methylbutanoate - Other names: 5-fluoro AMB and 5F-AMB.
24	[15]	methyl (1-(4-fluorobenzyl)-1H-indazole-3-carbonyl)-L-valinate2-
25		(1-(4-fluorobenzyl)-1H-indazole-3-carboxamido)-3-
26		methylbutanoate - Other names: FUB-AMB, MMB-FUBINACA,
27		and AMB-FUBINACA.
28	[16]	N-[1-(aminocarbonyl)-2,2-dimethylpropyl]-1-(cyclohexylmethyl)-1
29		H-indazole-3-carboxamide - Other names: MAB-CHMINACA and
30		ADB-CHMINACA.

1	[17]	Methyl 2-(1-(5-fluoropentyl)-1H-indazole-3-carboxamido)-3,3-
2		dimethylbutanoate - Other names: 5F-ADB and
3		5F-MDMB-PINACA.
4	[18]	N-(adamantan-1-yl)-1-(5-fluoropentyl)-1H-indazole-3-
5		carboxamide - Other names: 5F-APINACA and 5F-AKB48.
6	[19]	Methyl 2-(1-(cyclohexylmethyl)-1H-indole-3-carboxamido)-3,3-
7		dimethylbutanoate - Other names: MDMB-CHMICA and
8		MMB-CHMINACA.
9	[20]	Methyl 2-(1-(4-fluorobenzyl)-1H-indazole-3-carboxamido)-3,3-
10		dimethylbutanoate - Other names: MDMB-FUBINACA.
11	[21]	1-(4-cyanobutyl)-N-(2-phenylpropan-2-yl)-1H-indazole-3-carboxa
12		mide - Other names: 4-CN-CUMYL-BUTINACA; 4-cyano-
13		CUMYL-BUTINACA; 4-CN-CUMYL BINACA; CUMYL-4CN-
14		BINACA; SGT-78.
15	[22]	methyl 2-(1-(cyclohexylmethyl)-1H-indole-3-carboxamido)-
16		3-methylbutanoate - Other names: MMB-CHMICA, AMB-
17		CHMICA.
18	[23]	<u>1-(5-fluoropentyl)-N-(2-phenylpropan-2-yl)-1H-pyrrolo[2,3-b]pyridi</u>
19		ne-3-carboxamide - Other names: 5F-CUMYL-P7AICA.
20	(3) Indole car	boxylic acids. Any compound structurally derived from 1H-indole-
21	3-carboxy	lic acid or 1H-2-carboxylic acid substituted in both of the following
22	ways: at t	he nitrogen atom of the indole ring by an alkyl, haloalkyl,
23	cyanoalky	/l, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-2-
24	piperidiny	l)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-pyrrolidinyl)methyl,
25	1-(N-meth	nyl-3- morpholinyl)methyl, tetrahydropyranylmethyl, benzyl, or halo
26	benzyl gro	oup; and, at the hydroxyl group of the carboxylic acid by a phenyl,
27	benzyl, <u>cı</u>	<u>umyl,</u> naphthyl, adamantyl, cyclopropyl, or propionaldehyde group
28	whether c	or not the compound is further modified to any extent in the
29	following	ways:
30	(a) Sub	estitution to the indole ring to any extent; or

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

1		alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-
2		methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-(N-methyl-2-
3		pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl, or (tetrahydropyran-4-
4		yl)methyl group whether or not further substituted in the pyrrole ring to any
5		extent, whether or not substituted in the naphthyl ring to any extent.
6		Examples include: (5-(2-fluorophenyl)-1-pentylpyrrol-3-yl)-naphthalen-1-
7		ylmethanone - Other names: JWH-307.
8	(6)	Naphthylmethylindenes. Any compound containing a naphthylideneindene
9		structure with substitution at the 3-position of the indene ring by an alkyl,
10		haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl, cycloalkylethyl, 1-(N-methyl-
11		2-piperidinyl)methyl, 2 (4 morpholinyl)ethyl, 1-(N-methyl-2-
12		pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl, or (tetrahydropyran-4-
13		yl)methyl group whether or not further substituted in the indene ring to any
14		extent, whether or not substituted in the naphthyl ring to any extent.
15		Examples include: E-1-[1-(1-Naphthalenylmethylene)-1H-inden-3-yl]pentane
16		- Other names: JWH-176.
17	(7)	Cyclohexylphenols. Any compound containing a 2-(3-
18		hydroxycyclohexyl)phenol structure with substitution at the 5-position of the
19		phenolic ring by an alkyl, haloalkyl, cyanoalkyl, alkenyl, cycloalkylmethyl,
20		cycloalkylethyl, 1-(N-methyl-2-piperidinyl)methyl, 2-(4-morpholinyl)ethyl, 1-
21		(N-methyl-2-pyrrolidinyl)methyl, 1-(N-methyl-3-morpholinyl)methyl, or
22		(tetrahydropyran-4-yl)methyl group whether or not substituted in the
23		cyclohexyl ring to any extent. Examples include:
24		(a) 5-(1,1-dimethylheptyl)-2-[(1R,3S)-3-hydroxycyclohexyl]-phenol - Other
25		names: CP 47,497.
26		(b) 5-(1,1-dimethyloctyl)-2-[(1R,3S)-3-hydroxycyclohexyl]-phenol - Other
27		names: Cannabicyclohexanol and CP 47,497 C8 homologue.
28		(c) 5-(1,1-dimethylheptyl)-2-[(1R,2R)-5-hydroxy-2-(3-
29		hydroxypropyl)cyclohexyl]-phenol - Other names: CP 55,940.
30	(8)	Others specifically named:

	-		-	
1			(a)	(6aR,10aR)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan-2-yl)-
2				6a,7,10,10a-tetrahydrobenzo[c]chromen-1-ol - Other names: HU-210.
3			(b)	(6aS,10aS)-9-(hydroxymethyl)-6,6-dimethyl-3-(2-methyloctan-2-yl)-
4				6a,7,10,10a-tetrahydrobenzo[c]chromen-1-ol - Other names:
5				Dexanabinol and HU-211.
6			(C)	2,3-Dihydro-5-methyl-3-(4-morpholinylmethyl)pyrrolo[1,2,3-de]-1,4-
7				benzoxazin-6-yl]-1-napthalenylmethanone - Other names:
8				WIN 55,212-2.
9			(d)	Naphthalen-1-yl-(4-pentyloxynaphthalen-1-yl)methanone - Other
10				names: CB-13.
11	p.	Sub	stitute	d phenethylamines. This includes any compound, unless specifically
12		exc	epted,	specifically named in this schedule, or listed under a different
13		sch	edule,	structurally derived from phenylethan-2-amine by substitution on the
14		phe	nyl rin	g in any of the following ways, that is to say, by substitution with a fused
15		met	hylene	edioxy ring, fused furan ring, or fused tetrahydrofuran ring; by
16		sub	stitutic	on with two alkoxy groups; by substitution with one alkoxy and either
17		one	fused	furan, tetrahydrofuran, or tetrahydropyran ring system; or by
18		sub	stitutic	on with two fused ring systems from any combination of the furan,
19		tetra	ahydro	ofuran, or tetrahydropyran ring systems.
20		(1)	Whe	ther or not the compound is further modified in any of the following
21			ways	s, that is to say:
22			(a)	By substitution of phenyl ring by any halo, hydroxyl, alkyl,
23				trifluoromethyl, alkoxy, or alkylthio groups;
24			(b)	By substitution at the 2-position by any alkyl groups; or
25			(C)	By substitution at the 2-amino nitrogen atom with alkyl, dialkyl, benzyl,
26				hydroxybenzyl, methylenedioxybenzyl, or methoxybenzyl groups.
27		(2)	Exar	nples include:
28			(a)	2-(4-Chloro-2,5-dimethoxyphenyl)ethanamine (also known as 2C-C or
29				2,5-Dimethoxy-4-chlorophenethylamine).
30			(b)	2-(2,5-Dimethoxy-4-methylphenyl)ethanamine (also known as 2C-D or
31				2,5-Dimethoxy-4-methylphenethylamine).

1	(C)	2-(2,5-Dimethoxy-4-ethylphenyl)ethanamine (also known as 2C-E or
2		2,5-Dimethoxy-4-ethylphenethylamine).
3	(d)	2-(2,5-Dimethoxyphenyl)ethanamine (also known as 2C-H or 2,5-
4		Dimethoxyphenethylamine).
5	(e)	2-(4-lodo-2,5-dimethoxyphenyl)ethanamine (also known as 2C-l or
6		2,5-Dimethoxy-4-iodophenethylamine).
7	(f)	2-(2,5-Dimethoxy-4-nitro-phenyl)ethanamine (also known as 2C-N or
8		2,5-Dimethoxy-4-nitrophenethylamine).
9	(g)	2-(2,5-Dimethoxy-4-(n)-propylphenyl)ethanamine (also known as 2C-
10		P or 2,5-Dimethoxy-4-propylphenethylamine).
11	(h)	2-[4-(Ethylthio)-2,5-dimethoxyphenyl]ethanamine (also known as 2C-
12		T-2 or 2,5-Dimethoxy-4-ethylthiophenethylamine).
13	(i)	2-[4-(Isopropylthio)-2,5-dimethoxyphenyl]ethanamine (also known as
14		2C-T-4 or 2,5-Dimethoxy-4-isopropylthiophenethylamine).
15	(j)	2-(4-bromo-2,5-dimethoxyphenyl)ethanamine (also known as 2C-B or
16		2,5-Dimethoxy-4-bromophenethylamine).
17	(k)	2-(2,5-dimethoxy-4-(methylthio)phenyl)ethanamine (also known as
18		2C-T or 4-methylthio-2,5-dimethoxyphenethylamine).
19	(I)	1-(2,5-dimethoxy-4-iodophenyl)-propan-2-amine (also known as DOI
20		or 2,5-Dimethoxy-4-iodoamphetamine).
21	(m)	1-(4-Bromo-2,5-dimethoxyphenyl)-2-aminopropane (also known as
22		DOB or 2,5-Dimethoxy-4-bromoamphetamine).
23	(n)	1-(4-chloro-2,5-dimethoxy-phenyl)propan-2-amine (also known as
24		DOC or 2,5-Dimethoxy-4-chloroamphetamine).
25	(0)	2-(4-bromo-2,5-dimethoxyphenyl)-N-[(2-
26		methoxyphenyl)methyl]ethanamine (also known as 2C-B-NBOMe;
27		2,5B-NBOMe or 2,5-Dimethoxy-4-bromo-N-(2-
28		methoxybenzyl)phenethylamine).
29	(p)	2-(4-iodo-2,5-dimethoxyphenyl)-N-[(2 -
30		methoxyphenyl)methyl]ethanamine (also known as 2C-I-NBOMe; 2,5I-

1		NBOMe or 2,5-Dimethoxy-4-iodo-N-(2-
2		methoxybenzyl)phenethylamine).
3	(q)	N-(2-Methoxybenzyl)-2-(3,4,5-trimethoxyphenyl)ethanamine (also
4		known as mescaline-NBOMe or 3,4,5-trimethoxy-N-(2-
5		methoxybenzyl)phenethylamine).
6	(r)	2-(4-chloro-2,5-dimethoxyphenyl)-N-[(2-
7		methoxyphenyl)methyl]ethanamine (also known as 2C-C-NBOMe;
8		2,5C-NBOMe or 2,5-Dimethoxy-4-chloro-N-(2-
9		methoxybenzyl)phenethylamine).
10	(S)	2-(7-Bromo-5-methoxy-2,3-dihydro-1-benzofuran-4-yl)ethanamine
11		(also known as 2CB-5-hemiFLY).
12	(t)	2-(8-bromo-2,3,6,7-tetrahydrofuro [2,3-f][1]benzofuran-4-
13		yl)ethanamine (also known as 2C-B-FLY).
14	(u)	2-(10-Bromo-2,3,4,7,8,9-hexahydropyrano[2,3-g]chromen-5-
15		yl)ethanamine (also known as 2C-B-butterFLY).
16	(V)	N-(2-Methoxybenzyl)-1-(8-bromo-2,3,6,7-tetrahydrobenzo[1,2-b:4,5-
17		b']difuran-4-yl)-2-aminoethane (also known as 2C-B-FLY-NBOMe).
18	(w)	1-(4-Bromofuro[2,3-f][1]benzofuran-8-yl)propan-2-amine (also known
19		as bromo-benzodifuranyl-isopropylamine or bromo-dragonFLY).
20	(x)	N-(2-Hydroxybenzyl)-4-iodo-2,5-dimethoxyphenethylamine (also
21		known as 2C-I-NBOH or 2,5I-NBOH).
22	(y)	5-(2-Aminopropyl)benzofuran (also known as 5-APB).
23	(Z)	6-(2-Aminopropyl)benzofuran (also known as 6-APB).
24	(aa)	5-(2-Aminopropyl)-2,3-dihydrobenzofuran (also known as 5-APDB).
25	(bb)	6-(2-Aminopropyl)-2,3,-dihydrobenzofuran (also known as 6-APDB).
26	(cc)	2,5-dimethoxy-amphetamine (also known as 2,5-dimethoxy-a-
27		methylphenethylamine; 2,5-DMA).
28	(dd)	2,5-dimethoxy-4-ethylamphetamine (also known as DOET).
29	(ee)	2,5-dimethoxy-4-(n)-propylthiophenethylamine (also known as 2C-T-
30		7).
31	(ff)	5-methoxy-3,4-methylenedioxy-amphetamine.

1	(gg) 4-methyl-2,5-dimethoxy-amphetamine (also known as 4-methyl-2,5-	
2	dimethoxy-a-methylphenethylamine; DOM and STP).	
3	(hh) 3,4-methylenedioxy amphetamine (also known as MDA).	
4	(ii) 3,4-methylenedioxymethamphetamine (also known as MDMA).	
5	(jj) 3,4-methylenedioxy-N-ethylamphetamine (also known as N-ethyl-	
6	alpha-methyl-3,4(methylenedioxy)phenethylamine, MDE, MDEA).	
7	(kk) 3,4,5-trimethoxy amphetamine.	
8	(II) Mescaline (also known as 3,4,5-trimethoxyphenethylamine).	
9	q. Substituted tryptamines. This includes any compound, unless specifically	
10	excepted, specifically named in this schedule, or listed under a different	
11	schedule, structurally derived from 2-(1H-indol-3-yl)ethanamine (i.e., tryptamine	;)
12	by mono- or di-substitution of the amine nitrogen with alkyl or alkenyl groups or	
13	by inclusion of the amino nitrogen atom in a cyclic structure whether or not the	
14	compound is further substituted at the alpha-position with an alkyl group or	
15	whether or not further substituted on the indole ring to any extent with any alkyl,	,
16	alkoxy, halo, hydroxyl, or acetoxy groups. Examples include:	
17	(1) 5-methoxy-N,N-diallyltryptamine (also known as 5-MeO-DALT).	
18	(2) 4-acetoxy-N,N-dimethyltryptamine (also known as 4-AcO-DMT or O-	
19	Acetylpsilocin).	
20	(3) 4-hydroxy-N-methyl-N-ethyltryptamine (also known as 4-HO-MET).	
21	(4) 4-hydroxy-N,N-diisopropyltryptamine (also known as 4-HO-DIPT).	
22	(5) 5-methoxy-N-methyl-N-isopropyltryptamine (also known as 5-MeO-MiPT).	
23	(6) 5-methoxy-N,N-dimethyltryptamine (also known as 5-MeO-DMT).	
24	(7) Bufotenine (also known as 3-(Beta-Dimethyl-aminoethyl)-5-hydroxyindole;	
25	3-(2-dimethylaminoethyl)-5-indolol; N, N-dimethylserotonin; 5-hydroxy-N,N	I -
26	dimethyltryptamine; mappine).	
27	(8) 5-methoxy-N,N-diisopropyltryptamine (also known as 5-MeO-DiPT).	
28	(9) Diethyltryptamine (also known as N,N-Diethyltryptamine; DET).	
29	(10) Dimethyltryptamine (also known as DMT).	
30	(11) Psilocyn.	
31	r. 1-[3-(trifluoromethylphenyl)]piperazine (also known as TFMPP).	

1		S.	1-[4-(trifluoromethylphenyl)]piperazine.
2		t.	6,7-dihydro-5H-indeno-(5,6-d)-1,3-dioxol-6-amine (also known as 5,6-
3			Methylenedioxy-2-aminoindane or MDAI).
4	I	u.	2-(Ethylamino)-2-(3-methoxyphenyl)cyclohexanone (also known as
5			Methoxetamine or MXE).
6		V.	Ethylamine analog of phencyclidine (also known as N-ethyl-1-
7			phenylcyclohexylamine, (1-phenylcyclohexyl) ethylamine, N-(1-phenylcyclohexyl)
8			ethylamine, cyclohexamine, PCE).
9	١	W.	Pyrrolidine analog of phencyclidine (also known as 1-(1-phenylcyclohexyl)-
10			pyrrolidine, PCPy, PHP).
11		Х.	Thiophene analog of phencyclidine (also known as (1-[1-(2-thienyl) cyclohexyl]
12			piperidine; 2-Thienylanalog of phencyclidine; TPCP, TCP).
13		у.	1-[1-(2-thienyl)cyclohexyl]pyrrolidine (also known as TCPy).
14		Z.	Salvia divinorum, salvinorin A, or any of the active ingredients of salvia divinorum.
15	6.	Dep	ressants. Unless specifically excepted or unless listed in another schedule, any
16	I	mate	erial compound, mixture, or preparation which contains any quantity of the
17	1	follo	wing substances having a depressant effect on the central nervous system,
18	,	whe	never the existence of such salts, isomers, and salts of isomers is possible within
19	1	the	specific chemical designation:
20	;	a.	Flunitrazepam.
21	ł	b.	Gamma-hydroxybutyric acid.
22	C.	<u>b.</u>	Mecloqualone.
23	d.	<u>C.</u>	Methaqualone.
24	7. 3	Stim	ulants. Unless specifically excepted or unless listed in another schedule, any
25	I	mate	erial, compound, mixture, or preparation which contains any quantity of the
26	1	follo	wing substances having a stimulant effect on the central nervous system,
27	i	inclu	uding its salts, isomers, and salts of isomers:
28	;	a.	Aminorex (also known as 2-amino-5-phenyl-2-oxazoline, or 4,5-dihydro-5-phenyl-
29			2-oxazolamine).
30	I	b.	Cathinone.

1	C.	Sub	stitute	d cathinones. Any compound, material, mixture, preparation, or other
2				nless listed in another schedule or an approved food and drug
3		•		ation drug (e.g., buproprion, pyrovalerone), structurally derived from 2-
4				pan-1-one by substitution at the 1-position with either phenyl, naphthyl,
5				ne ring systems, whether or not the compound is further modified in
6			•	following ways:
7		(1)		ubstitution in the ring system to any extent with alkyl, alkylenedioxy,
8		()	•	ky, haloalkyl, hydroxyl, or halide substituents, whether or not further
9				tituted in the ring system by one or more other univalent substitutents;
10		(2)		ubstitution at the 3-position with an acyclic alkyl substituent;
11		(3)	-	ubstitution at the 2-amino nitrogen atom with alkyl, dialkyl, benzyl, or
12		. ,	meth	noxybenzyl groups; or
13		(4)	By in	iclusion of the 2-amino nitrogen atom in a cyclic structure.
14			Som	e trade or other names:
15			(a)	3,4-Methylenedioxy-alpha-pyrrolidinopropiophenone (also known as
16				MDPPP).
17			(b)	3,4-Methylenedioxy-N-ethylcathinone (also known as Ethylone,
18				MDEC, or bk-MDEA).
19			(C)	3,4-Methylenedioxy-N-methylcathinone (also known as Methylone or
20				bk-MDMA).
21			(d)	3,4-Methylenedioxypyrovalerone (also known as MDPV).
22			(e)	3,4-Dimethylmethcathinone (also known as 3,4-DMMC).
23			(f)	2-(methylamino)-1-phenylpentan-1-one (also known as Pentedrone).
24			(g)	2-Fluoromethcathinone (also known as 2-FMC).
25			(h)	3-Fluoromethcathinone (also known as 3-FMC).
26			(i)	4-Methylethcathinone (also known as 4-MEC and 4-methyl-N-
27				ethylcathinone).
28			(j)	4-Fluoromethcathinone (also known as Flephedrone and 4-FMC).
29			(k)	4-Methoxy-alpha-pyrrolidinopropiophenone (also known as MOPPP).
30			(I)	4-Methoxymethcathinone (also known as Methedrone; bk-PMMA).
31			(m)	4'-Methyl-alpha-pyrrolidinobutiophenone (also known as MPBP).

	-	-	
1		(n)	Alpha-methylamino-butyrophenone (also known as Buphedrone or
2			MABP).
3		(0)	Alpha-pyrrolidinobutiophenone (also known as alpha-PBP).
4		(p)	Alpha-pyrrolidinopropiophenone (also known as alpha-PPP).
5		(q)	Alpha-pyrrolidinopentiophenone (also known as Alpha-
6			pyrrolidinovalerophenone or alpha-PVP).
7		(r)	Beta-keto-N-methylbenzodioxolylbutanamine (also known as Butylone
8			or bk-MBDB).
9		(s)	Ethcathinone (also known as N-Ethylcathinone).
10		(t)	4-Methylmethcathinone (also known as Mephedrone or 4-MMC).
11		(u)	Methcathinone.
12		(v)	N,N-dimethylcathinone (also known as metamfepramone).
13		(w)	Naphthylpyrovalerone (naphyrone).
14		(x)	B-Keto-Methylbenzodioxolylpentanamine (also known as Pentylone).
15		(y)	4-Methyl-alpha-pyrrolidinopropiophenone (also known as 4-MePPP
16			and MPPP).
17		<u>(z)</u>	1-(1,3-benzodioxol-5-yl)-2-(ethylamino)-pentan-1-one (also known as
18			Ephylone and N-Ethylpentylone).
19	d.	Fenethylli	ne.
20	e.	Fluoroam	phetamine.
21	f.	Fluorome	thamphetamine.
22	g.	(±)cis-4-m	nethylaminorex (also known as (±)cis-4,5-dihydro-4-methyl-5-phenyl-2-
23		oxazolam	ine).
24	h.	N-Benzyl	piperazine (also known as BZP, 1-benzylpiperazine).
25	i.	N-ethylan	nphetamine.
26	j.	N, N-dime	ethylamphetamine (also known as N,N-alpha-trimethyl-
27		benzenee	thanamine; N,N-alpha-trimethylphenethylamine).
28	SECTIC	ON 3. AMEN	DMENT. Subsection 7 of section 19-03.1-07 of the North Dakota
29	Century Coo	de is amend	ed and reenacted as follows:
30	7. Ha	Illucinogenic	substances.

1	<u>a</u> .	Nabilone [another name for nabilone (±)-trans-3-(1, 1-dimethylheptyl)-6, 6a, 7, 8,
2		10, 10a-hexahydro-1-hydroxy-6, 6-dimethyl-9Hdibenzo [b, d] pyran-9-one].
3	<u>b</u> .	Dronabinol [(-)-delta-9-trans tetrahydrocannabinol] in an oral solution in a drug
4		product approved for marketing by the federal food and drug administration.
5	SECTI	ON 4. AMENDMENT. Subsection 4 of section 19-03.1-09 of the North Dakota
6	Century Co	de is amended and reenacted as follows:
7	4. D	epressants. Unless specifically excepted or unless listed in another schedule, any
8	m	aterial, compound, mixture, or preparation that contains any quantity of the following
9	SI	ibstances having a depressant effect on the central nervous system:
10	a	Any compound, mixture, or preparation containing:
11		(1) Amobarbital;
12		(2) Secobarbital;
13		(3) Pentobarbital;
14		or any salt thereof and one or more other active medicinal ingredients which are
15		not listed in any schedule.
16	b	Any suppository dosage form containing:
17		(1) Amobarbital;
18		(2) Secobarbital;
19		(3) Pentobarbital;
20		or any salt of any of these drugs and approved by the food and drug
21		administration for marketing only as a suppository.
22	C	Any substance that contains any quantity of a derivative of barbituric acid, or any
23		salt of a derivative of barbituric acid, except those substances which are
24		specifically listed in other schedules thereof.
25	d	Chlorhexadol.
26	e	Embutramide.
27	f.	Gamma-hydroxybutyric acid in a United States food and drug administration-
28		approved drug product.
29	g	Ketamine.
30	h	Lysergic acid.
31	i.	Lysergic acid amide.

1	j.	Methyprylon.		
2	k.	Perampanel.		
3	I.	Sativex or its successor name as determined by the federal food and drug		
4		administration.		
5	<u>m.</u>	Sulfondiethylmethane.		
6	m.<u>n.</u>	Sulfonethylmethane.		
7	n.<u>o.</u>	Sulfonmethane.		
8	0. p.	Tiletamine and zolazepam or any salt thereof. Some trade or other names for a		
9		tiletamine-zolazepam combination product: Telazol. Some trade or other names		
10		for tiletamine: 2-(ethylamino)-2-(2-thienyl)-cyclohexanone. Some trade or other		
11		names for zolazepam: 4-2(2-fluorophenyl)-6, 8-dihydro-1,3,8-trimethylpyrazolo-		
12		[3,4-e][1,4]-diazepin-7(1H)-one, flupyrazapon.		
13	SECTION	N 5. AMENDMENT. Subsection 7 of section 19-03.1-11 of the North Dakota		
14	Century Code	e is amended and reenacted as follows:		
15	7. Oth	er substances. Unless specifically excepted or unless listed in another schedule,		
16	any	material, compound, mixture, or preparation which contains any quantity of:		
17	a.	Pentazocine, including its salts.		
18	b.	Butorphanol, including its optical isomers.		
19	С.	Eluxadoline (5-[[[(2S)-2-amino-3-[4-aminocarbonyl)-2,6-dimethylphenyl]-1-		
20		oxopropyl][(1S)-1-(4-phenyl-1H-imidazol-2-yl)ethyl]amino]methyl]-2-		
21		methoxybenzoic acid) (including its optical isomers) and its salts, isomers, and		
22		salts of isomers.		
23	d.	Epidiolex or its successor name as determined by the United States food and		
24		drug administration.		
25	SECTION	N 6. AMENDMENT. Subsection 5 of section 19-03.1-13 of the North Dakota		
26	Century Code	e is amended and reenacted as follows:		
27	5. Dep	pressants. Unless specifically exempted or excluded or unless listed in another		
28	sch	edule, any material, compound, mixture, or preparation that contains any quantity		
29	of th	the following substances having a depressant effect on the central nervous system,		
30	inclu	uding its salts, isomers, and salts of isomers whenever the existence of such salts,		
31	ison	ners, and salts of isomers is possible:		

1	a.	Brivaracetam ((2S)-2-[(4R)-2-oxo-4-propylpyrrolidin-1-yl]butanamide) (also referred
2		to as BRV; UCB-34714; Briviact) (including its salts).
3	b.	Ezogabine N-[2-amino-4-(4-fluorobenzylamino)-phenyl]-carbamic acid ethyl ester.
4	C.	Lacosamide [(R)-2-acetoamido-N-benzyl-3-methoxy-propionamide].
5	d.	Pregabalin [(S)-3-(aminomethyl)-5-methylhexanoic acid].
6	<u>e.</u>	Approved cannabidiol drugs. A drug product in finished dosage formulation that
7		has been approved by the federal food and drug administration, which contains
8		cannabidiol (2-[1R-3-methyl-6R-(1-methylethenyl)-2-cyclohexen-1-yl]-5-pentyl-
9		1,3-benzenediol) derived from cannabis and no more than 0.1 percent weight for
10		weight residual tetrahydrocannabinols.
11	<u>f.</u>	Gabapentin [2-[1-(aminomethyl) cyclohexyl] acetic acid].
12	SECTION	7. EMERGENCY. This Act is declared to be an emergency measure.