



Restrictive Practices Reporting Form

Important information

This form is approved by the NDIS Quality and Safeguards Commissioner for the purposes of section 14 of the *National Disability Insurance Scheme (Restrictive Practices and Behaviour Support) Rules 2018*.

Privacy

This form seeks to collect information—including personal information—for the purpose of administering and enforcing the *National Disability Insurance Scheme Act 2013* and *National Disability Insurance Scheme (Restrictive Practices and Behaviour Support) Rules 2018*. Please refer to the Privacy Collection Statement and the NDIS Commission's Privacy Policy at www.ndiscommission.gov.au

Security

Once the NDIS Quality and Safeguards Commission (NDIS Commission) receives information from you via email or any other means, the information is in a secure environment. Your personal information will not be released unless the law permits it or your permission is granted.

You need to be aware of inherent risks associated with the transmission of information via email and otherwise over the internet. If you have concerns in this regard, the NDIS Commission has other ways of obtaining and providing information including mail, telephone and FilePoint. For advice about how to use FilePoint, please contact the NDIS Commission at 1800 035 544.

Instructions

This form must be completed by registered NDIS providers in New South Wales and South Australia as required under the *National Disability Insurance Scheme (Restrictive Practices and Behaviour Support) Rules 2018*. Reporting is required from providers that use regulated restrictive practices.

The requirement to report to the NDIS Commission does not replace existing obligations on providers to report to other relevant authorities, including child protection agencies or police.

Once completed, email the form together with relevant documents to behavioursupport@ndiscommission.gov.au

When completed, this document contains information submitted to the NDIS Quality and Safeguards Commission (the NDIS Commission) by a third party for the purposes of the *National Disability Insurance Scheme Act 2013* (Cth). The NDIS Commission makes no representations about, and accepts no liability for, the accuracy of information in this document.

Provider details

Organisation _____

Outlet _____

Authorised Reporting Officer _____

Phone _____

Email _____

Person details

Name _____

Date of birth _____

Address _____

Type of residence _____

Phone number _____

Email _____

Preferred method of contact _____

NDIS participant number _____

Guardian appointed _____

Guardian functions _____

Consent to share information received? _____

Consent received from _____

Date consent received _____

Behaviour support planning status _____

Behaviour support plan ID (provided by NDIS Commission once submitted) _____

Jurisdiction (State) _____

Reporting of Regulated Restrictive Practices

Reporting month (use calendar months only) _____

*note if fortnightly reporting is required where State or Territory short term authorisation is issued indicate reporting period

Start _____ Finish _____

Routine Use — where details of the practice are contained in the behaviour support plan

Regulated Restrictive Practice (RRP)	Restrictive practice sub-type	Authorisation and consent received	Used in accordance with plan (yes/no)	Variation from plan — describe any variation from the plan if occurred including how and why
E.g. Chemical		Yes	Yes	Nil

PRN Use — where details of the practice are contained in the behaviour support plan

For PRN use, details of each instance of the restrictive practice must be recorded.

Restrictive practice use details

Restrictive practice type _____

Sub-type _____

Administration type _____

Describe how the practice was used (if chemical restraint, provide medication details further down under “Chemical restraint — medication details”)

Date used _____

Start time _____

Finish time _____

Duration _____

Where was the practice used? (e.g. living room, at the shops in the community)

Names and contact details of persons who administered the RP

Names and contact details of any witnesses

Describe the impact on the person from the use of the practice

Was there any injury to the person? If yes, please provide details

Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

Restrictive practice type _____

Sub-type _____

Administration type _____

Describe how the practice was used (if chemical restraint, provide medication details further down under “Chemical restraint — medication details”)

Date used _____

Start time _____

Finish time _____

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Where was the practice used? (e.g. living room, at the shops in the community)

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Describe the impact on the person from the use of the practice

Was there any injury to the person? If yes, please provide details

Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

Restrictive practice type _____

Sub-type _____

Administration type _____

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Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

Restrictive practice type _____

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Administration type _____

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Date used _____

Start time _____

Finish time _____

Duration _____

Where was the practice used? (e.g. living room, at the shops in the community)

Names and contact details of persons who administered the RP

Names and contact details of any witnesses

Describe the impact on the person from the use of the practice

Was there any injury to the person? If yes, please provide details

Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

(If additional fields are required include additional information as an attachment)

Chemical restraint — medication details

If medication details are not included in the behaviour support plan, please complete details below

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 1 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects _____

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 2 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects _____

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 3 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects _____

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 4 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects _____

Reporting of Regulated Restricted Practices not detailed in a behaviour support plan but authorisation from the state body

Please provide full details of any Regulated Restrictive Practices that are being used and are not currently detailed in a current behaviour support plan.

For PRN use, details of each instance of the restrictive practice must be recorded.

For routine use, indicate the pattern of use

Restrictive practice 1

Restrictive practice type _____

Sub-type _____

Administration type _____

Describe how the practice was used (if chemical restraint, provide medication details further down under “Chemical restraint — medication details”)

Date used _____

Start time _____

Finish time _____

Duration _____

Where was the practice used? (e.g. living room, at the shops in the community)

Names and contact details of persons who administered the RP

Names and contact details of any witnesses

Describe the impact on the person from the use of the practice

Was there any injury to the person? If yes, please provide details

Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

(If additional fields are required include additional information as an attachment)

Restrictive practice 2

Restrictive practice type _____

Sub-type _____

Administration type _____

Describe how the practice was used (if chemical restraint, provide medication details further down under “Chemical restraint — medication details”)

Date used _____

Start time _____

Finish time _____

Duration _____

Where was the practice used? (e.g. living room, at the shops in the community)

Names and contact details of persons who administered the RP

Names and contact details of any witnesses

Describe the impact on the person from the use of the practice

Was there any injury to the person? If yes, please provide details

Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

(If additional fields are required include additional information as an attachment)

Restrictive practice 3

Restrictive practice type _____

Sub-type _____

Administration type _____

Describe how the practice was used (if chemical restraint, provide medication details further down under “Chemical restraint — medication details”)

Date used _____

Start time _____

Finish time _____

Duration _____

Where was the practice used? (e.g. living room, at the shops in the community)

Names and contact details of persons who administered the RP

Names and contact details of any witnesses

Describe the impact on the person from the use of the practice

Was there any injury to the person? If yes, please provide details

Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

(If additional fields are required include additional information as an attachment)

Restrictive practice 4

Restrictive practice type _____

Sub-type _____

Administration type _____

Describe how the practice was used (if chemical restraint, provide medication details further down under “Chemical restraint — medication details”)

Date used _____

Start time _____

Finish time _____

Duration _____

Where was the practice used? (e.g. living room, at the shops in the community)

Names and contact details of persons who administered the RP

Names and contact details of any witnesses

Describe the impact on the person from the use of the practice

Was there any injury to the person? If yes, please provide details

Was the use of the practice a reportable incident? _____

Describe why the regulated restrictive practice was used

Description of behaviour of the person that lead to the use of the regulated restrictive practice

What actions taken in response to the use of the practice?

What less restrictive options were considered prior to the use of the practice?

(If additional fields are required include additional information as an attachment)

Chemical restraint — medication details

If medication details are not included in the behaviour support plan, please complete details below

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 1 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects _____

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 2 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects _____

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 3 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects

Important note about chemical restraints

All medication must be administered according to a person's current medical record only; information in the behaviour support plan is to record details of chemical restraint for data collection purposes only and should not be used for administering any medication.

Medication 4 details

Drug name (Refer to Attachment A for a list of chemical restraint drugs)

Administration type _____

Dosage _____

Unit of measurement _____

Frequency _____

Unit of measurement (day, week, month) _____

Route _____

Prescriber _____

Name _____

Date of last review by Doctor _____

Conditions/limits of use _____

Side effects _____

Declaration

I declare that:

- I am duly authorised by the provider in this form to submit this regulated restrictive practices report.
- I understand that this information is being collected by the NDIS Quality and Safeguards Commission (NDIS Commission) for the purposes outlined in *the NDIS (Restrictive Practices and Behaviour Support) Rules 2018*.
- I have read the NDIS Commission's NDIS restrictive practices and behaviour support guidance and understand the requirements of registered NDIS Providers in relation to notifying the NDIS Commission of the use of regulated restrictive practices.
- I understand that the NDIS Commission will, if required, use the information to undertake compliance and enforcement activities consistent with *the National Disability Insurance Scheme Act 2013 (the Act)* and any Rules established under the Act.
- I acknowledge the NDIS Commission may share the information contained in the application form with relevant Commonwealth, state and territory agencies, including the police.
- To the best of my knowledge, the information provided in this application is true, correct and accurate.
- I acknowledge that the giving of false or misleading information to the Commonwealth is a serious offence under section 137.1 of the schedule to the *Criminal Code Act 1995*.

Signature _____

Full name _____

Date _____

Job title _____

Please email completed form with all attachments to:
behavioursupport@ndiscommission.gov.au

Office use only

Plan ID number _____

Date form received _____

Date entered in CBAS _____

Entered by _____

Attachment A: Chemical restraint drug names

Abilify	Aromasin	Celapram	Clomipramine hydrochloride
Agrylin	Aropax	Celica	Clonazepam
Akineton	Artane	Certirizine	Clopine
Aldazine	Ativan	Cetuximab (rmc)	Clopixol
Alemtuzumab	Atomoxetine hydrochloride	Chemmart Alprazolam	CloSyn
Alepam	Attenta	Chemmart Citalopram	Clozapine
Allegron	Aurorix	Chemmart Clomipramine	Clozaril
Alodorm	Auscap 20 mg Capsules	Chemmart Diazepam	Cogentin
Alprax	Avanza	Chemmart Fluoxetine	Comtan
Alprazolam	Avanza SolTab	Chemmart Gabapentin	Concerta Extended-Release Tblt
Alprazolam-DP	Avastin	Chemmart Methylphenidate	Concorz
Amantadine hydrochloride	Axit	Chemmart Mirtazapine	Copper, intrauterine device
Aminoglutethimide	Benzhexol hydrochloride	Chemmart Moclobemide	Cosudex 150 mg
Amira	Benztrop	Chemmart Paroxetine	Cosudex 50 mg
Amisulpride	Benztropine mesylate	Chemmart Sertraline	Cymbalta
Amisulpride Sandoz	Bevacizumab (rch)	Chemmart Tamoxifen	Cyprohexal 100 mg
Amisulpride Winthrop	Bicalutamide	Chloral hydrate	Cyprohexal 50 mg
Amitriptyline hydrochloride	Biperiden hydrochloride	Chloral Hydrate Mix 1 g/10 mL	Cyprone
Anafranil	Bortezomib	Chlorpromazine hydrochloride	Cyprostat 50 mg
Anagrelide hydrochloride	Brenda-35 ED	Chlorpromazine hydrochloride	Cyprostat-100
Anandron	Brevinor, Brevinor-1, Synphasic	Chlorpromazine Mixture	Cyproterone acetate
Anastrozole	Bromazepam	Ciazil	Cytadren
Anatensol	Buspar	Cipramil	Dacarbazine
Androcur	Buspirone hydrochloride	Cisplatin	Dacarbazine for Injection(DBL)
Androcur-100	Cabaser	Cisplatin Ebewe	Dacarbazine Sandoz
Anexate	Cabergoline	Cisplatin Injection	Dasatinib
Antenex	Camptosar	Cisplatin Injection (DBL)	DBL Gabapentin Capsules
APO-go	Carbamazepine	Citalobell	Depo-Provera
Apomine Injection	Carbamazepine Sandoz	Citalopram 20	Depo-Ralovera
Apomorphine hydrochloride	Carbamazepine-BC	Citalopram hydrobromide	Deptran
Aricept	Carboplatin	Citalopram Winthrop	Dexamphetamine sulfate
Arima	Carboplatin Ebewe	Citalopram-RL	Dexamphetamine Tablets
Arimidex	Carboplatin Injection	Clobazam	
Aripiprazole	Carboplatin Injection (DBL)	Clobemix	

Dexmedetomidine hydrochloride	Erbitux 5 mg/mL	Flutamide	Haloperidol decanoate
Diane-35 ED	Erlotinib hydrochloride	Flutamin	Herceptin
Diazepam	Escitalopram oxalate	Fluvoxamine maleate	Hycamtin
Diazepam Elixir 10 mg/10 mL	Esipram	Frisium	Hypnodorm
Diazepam Injection (DBL)	Estelle-35 ED	Fulvestrant	Hypnovel
Diazepam-DP	Ethinylloestradiol	Gabahexal	Imatinib mesylate
Dilantin	Ethosuximide	Gabapentin	Imipramine hydrochloride
Donepezil hydrochloride	Etonogestrel	Gabapentin 300, 400	Imovane
Dormizol	Etopophos	Gabaran	Implanon Implant
Dosulepin	Etoposide	Gabitril	Imrest
Dothep	Etoposide Ebewe	Galantamine hydrobromide	Invega
Dothiepin hydrochloride	Etoposide Injection	Gantin	Iressa
Douglas Gabapentin Capsules	Etoposide Injection (DBL)	Gefitinib	Irinotecan hydrochloride
Douglas-Methylphenidate	Etoposide phosphate	Genox	Irinotecan Injection Concnrte
Doxepin hydrochloride	Eulexin	GenRx Alprazolam	Irinotecan Sandoz
Doxylamine succinate	Exelon	GenRx Citalopram	Juliet-35 ED
Dozile	Exemestane	GenRx Clomipramine	Kalma
Droleptan Injection	Extine	GenRx Cyproterone Acetate 50mg	Keppra
Droperidol	Fareston	GenRx Cyproterone Acetate100mg	Kinson
Ducene	Faslodex	GenRx Diazepam	Lamictal
Duloxetine hydrochloride	Faverin	GenRx Fluoxetine	Lamidus
Ebixa	Femara	GenRx Gabapentin Capsules	Lamitrin
Edronax	Femoden ED	GenRx Lamotrigine	Lamogine
Efexor	Fluanxol	GenRx Methylphenidate	Lamotrigine generichealth
Efexor-XR	Flumazenil	GenRx Mirtazapine	Lamotrigine-DP
Eldepryl	Flumazenil Injection DBL	GenRx Moclobemide	Lapatinib ditosylate
Eleva	Flunitrazepam	GenRx Paroxetine	Largactil
Eligard	Fluohexal	GenRx Sertraline	Letrozole
Elmendos	Fluoxetine 20	GenRx Tamoxifen	Leuprorelin acetate
Eloxatin	Fluoxetine hydrochloride	GenRx Zolpidem	Levetiracetam
Endep	Fluoxetine-DP	Glivec	Levlen ED
Entacapone	Flupenthixol decanoate	Goserelin acetate	Levodopa
Epilim	Fluphenazine decanoate	Halcion	Levohexal
Erbitux	Fluphenazine hydrochloride	Haldol Decanoate	Levonelle-2
		Haloperidol	

Levonorgestrel	Microgynon 30 ED	Nordette	Pericyazine
Lexapro	Microgynon 50 ED	Norethisterone	Permax
Lexotan	Microlevlen ED	Noriday 28	Pharmacor Gabapentin
Lithicarb	Microlut	Norimin, Norimin-1, Improvil 28Day	Phenelzine sulfate
Lithium carbonate	Micronor	Norinyl-1	Phenobarbitone
Locilan 28 Day	Microval	NorLevo	Phenobarbitone Elixir 15mg/5mL
Loette	Midazolam hydrochloride	Normison	Phenobarbitone Injection (DBL)
Logynon ED	Midazolam Injection	Nortriptyline hydrochloride	Phenobarbitone sodium
Loratidine	Midazolam Injection BP (DBL)	Nupentin	Phenobarbitone Sodium Inj/Tblt
Lorazepam	Midazolam Sandoz	NuvaRing	Phenytoin Injection BP (DBL)
Lovan	Minulet	Oestrogens, conjugated	Phenytoin sodium
Lucrin	Mirena	Olanzapine	Pimozide
Lucrin Depot	Mirtazapine	Orap	Placil
Lumin	Mirtazapine Sandoz	Ortho-Gynol	Postinor-2
Luvox	Mirtazapine-DP	Ospolot	Pramipexole hydrochloride
Lyrica	Mirtazon	Oxalatin	Precedex
MabCampath	Moclobemide	Oxaliplan	Pregabalin
Mabthera	Modafinil	Oxaliplatin	Premarin Tablets
Madopar	Modavigil	Oxaliplatin (DBL)	Premia (2.5/5 Continuous)
Maosig	Modecate	Oxaliplatin Ebewe	Primidone
Marvelon 28	Mogadon	Oxazepam	Primolut N
Medroxyhexal	Mohexal	Oxcarbazepine	Pristiq
Medroxyprogesterone acetate	Monofeme	Oxetine	Procarbazine hydrochloride
Megace	Movox	Paliperidone	Procur
Megestrol acetate	Multiload-cu 250, cu 375/ SL	Paraldehyde	Procur 100
Melatonin	Murelax	Paraldehyde Injection BP (DBL)	Promethazine (Phenergan)
Melatonin	Mysoline	Parnate	Prothiaden
Memantine hydrochloride	Nardil	Paroxetine 20	Provera
Mestranol	Natulan	Paroxetine hydrochloride	Prozac
Methyl amino hydrochloride	Neulactil	Paroxetine Winthrop	Quetiapine fumarate
Methylphenidate hydrochloride	Neurontin	Paroxetine-DP	Quilonum SR
Metvix	Nexavar	Paxam	Ralovera
Mianserin hydrochloride	Nilotinib	Paxtine	
Microgynon 20 ED	Nilutamide	Pendine	
Microgynon 30	Nitrazepam	Pergolide mesylate	
	Nolvadex, Nolvadex-D		

Reboxetine mesylate	Sodium Valproate Sandoz	Fluoxetine Terry White Chem	Unisom Sleepgels
Remeron	Solian Tablets and Solution	Gabapentin Terry White Chem	Valette
Reminyl	Somidem	Methylphenidol Terry White Chem	Valium
Restavit Tablets	Sorafenib tosylate	Mirtazapine Terry White Chem	Valpam
Rilutek	Sprycel	Moclobemide Terry White Chem	Valpro
Riluzole	Stalevo	Paroxetine Terry White Chem	Valproate Winthrop
Risperdal	Stelazine	Sertraline Terry White Chem	Velcade
Risperdal Consta	Stildem	Diazepam Terry White Chemists	Venlafaxine hydrochloride
Risperdal Consta	Stilnox	Tamoxifen Terry White Chemists	Vepesid
Risperidone	Stilnox CR	Tetrabenazine	Vesanoid
Risperidone Quicklet	Strattera	Tetrabenazine	Vigabatrin
Ritalin 10, Ritalin LA	Sulthiame	Thioridazine hydrochloride	Vimpat
Rituximab	Sunitinib malate	Tiagabine hydrochloride	Voxam
Rivastigmine hydrogen tartrate	Surmontil	Tofranil	Vumon
Rivotril	Sutent	Tolerade	Winthrop Oxaliplatin
Rixadone	Symmetrel	Tolvon	Xanax
Sabril	Talam	Topamax	Xydep
Seaze	Talohexal	Topiramate	Yasmin
Selegiline hydrochloride	Tamosin	Topotecan hydrochloride	Zactin
Selgene	Tamoxen	Toremifene citrate	Zamhexal
Serenace	Tamoxifen citrate	Tranylcypromine sulfate	Zarontin
Serepax	Tamoxifen Hexal	Trastuzumab	Zeldox
Seroquel	Tarceva	Triazolam	Ziprasidone hydrochloride
Sertra	Tasigna	Trifeme	ZolaCos CP
Sertraline 50, 100	Tegretol	Trifluoperazine hydrochloride	Zoladex 10.8 mg Implant
Sertraline generichealth	Temaze	Trileptal	Zoladex 3.6 mg Implant
Sertraline hydrochloride	Temazepam	Trimipramine	Zoloft
Sertraline Winthrop	Temtabs	Triphasil	Zolpidem Sandoz
Sertraline-DP	Teniposide	Triquilar ED	Zolpidem tartrate
Setrona	Teril	Tykerb	Zolpidem-DP
Sifrol	Terry White Chem		Zopiclone
Sinemet	Alprazolam		Zuclopenthixol
Sinemet CR	Terry White Chem		Zuclopenthixol acetate
Sinequan	Citalopram		Zyprexa
Snuzaid Gels	Terry White Chem		Zyprexa IM
Snuzaid Tabs	Clomipramine		
Sodium valproate	Terry White Chem		