GOVERNMENT OF GOA, DAMAN AND DIU
Public Health Department

Notification
13/53/87-I/PHD

In exercise of the powers conferred by section 78 read with section 10, sub-section (2) of section 71 and section 65 of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and of all other powers enabling it in this behalf, the Government of Goa, Daman and Diu hereby makes the following rules, namely:

1. Short title, extent and commencement. — (1) These rules may be called the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987.

(2) They extend to the whole of the Union territory of Goa, Daman and Diu.

(3) They shall come into force at once.

2. Definitions. — In these rules, unless there is anything repugnant in the subject or context,

(a) “Act” means the Narcotic Drugs and Psychotropic Substances Act, 1985 (61 of 1985);

(b) “authorised Officer” means,

(i) in respect of Chapter I, II and III, the “Commissioner of Excise” of Goa, Daman and Diu and any Officer authorised by him in this behalf by notification in the Official Gazette;

(ii) in respect of chapter IV and V, the Drugs Controller, Directorate of Health Services, Panaji, Goa or any other officer authorised by him in this behalf by notification in the Official Gazette;

(c) “Chemical Analyser” means the Chief Medical Officer, Public Health Laboratory, Directorate of Health Services, Panaji-Goa, and includes any officer authorised by him in this behalf or any officer not below the rank of Junior Scientific Officer attached to the Office of Drugs Controller, Directorate of Health Services, Panaji-Goa, and appointed for this purpose;

(d) “Drugs Controller” means the Drugs Controller, Office of the Drugs Controller, Government of Goa, Daman and Diu, appointed under the provisions of rule 50 of the Drugs and Cosmetics Rules, 1945;

(e) “Commissioner of Excise” means an Officer appointed under sub-section (1) of section 3 of the Goa, Daman and Diu Excise Duty Act, 1964 (Act 5 of 1964);

(f) “Form” means the form appended to these rules;

(g) “Government” means the Government of Goa, Daman and Diu;

(h) “Inspector” means an Inspector appointed under section 21 of the Drugs and Cosmetics Act, 1940 (Central Act 23 of 1940);

(i) “Licence” means a licence granted under these rules;

(j) “Licensing Authority” means any Officer of the Drugs Controller’s Office, not below the rank of the Assistant Drugs Controller, authorised or designated as such by the Drugs Controller by an order published in the Official Gazette.

(k) “Licensed Chemist” means a person licensed under these rules for the possession and sale or dispensing of any prescription of manufactured drugs and any preparation containing any manufactured drug;

(l) “Licensed Dealer in manufactured drugs” means a person who has obtained a licence under these rules for the manufacture, possession and sale of any preparation containing any manufactured drug from the material which the maker is lawfully entitled to possess;

(m) “Medical Board” means a Board constituted under these rules;
(n) "Medical Practitioner" means a person —
(i) holding a qualification granted by an authority specified or notified under section 3 of the Indian Medical Degrees Act, 1916 (Central Act 7 of 1916), or specified in the Schedules to the Indian Medical Council Act, 1956 (Central Act 102 of 1956); or
(ii) registered or eligible for registration in a medical register of union territory meant for the registration of persons practising the modern scientific system of medicine; or
(iii) registered in a medical register of union territory, who although not falling within sub-clause (i) or sub-clause (ii) is declared by a general or special order made by the Government, in this behalf as a person practising the modern scientific system of medicine for the purpose of this Act; or
(iv) registered or eligible for registration in the register of dentists for a Union territory under the Dentists Act, 1948 (Central Act 16 of 1948); or
(v) who is engaged in the practice of veterinary medicine and who possesses qualifications approved by the Government;
(o) "Pass" means a pass granted under these rules;
(n) "Permit" means a permit granted under these rules;
(q) "Union territory of Goa, Daman and Diu" means the Union territory of the Goa, Daman and Diu.

CHAPTER I

Poppy straw

3. Prohibition of possession, sale, etc of poppy straw: — Possession, sale, import inter-state, export inter-state, transport, export from India, consumption and use of poppy straw is prohibited.

CHAPTER II

Opium

4. Licence for possessing opium by registered medical Practitioner: — (1) Any registered medical practitioner desiring to possess opium for use as an ingredient in any medicine and to sell such medicines containing opium shall make an application to the authorised officer for a licence in that behalf.

(2) On receipt of an application under sub-rule (1), the authorised officer shall make such inquiries as he deems necessary and if he is satisfied that there is no objection to grant the licence applied for, he may, subject to the orders of Commissioner of Excise, if any, grant the applicant a licence in 'Form — O.P. I' on payment of a fee of rupees fifty only.

6. Licence for possessing opium by dealer. — (1) Any person desiring to possess and sell medicines containing opium shall make an application to the authorised officer for a licence in that behalf.

(2) On receipt of an application under sub-rule (1), the authorised officer shall make such inquiries as he deems necessary and if he is satisfied that there is no objection to grant the licence applied for, he may, subject to the orders of Commissioner of Excise, if any, grant the applicant a licence in 'Form — O.P. II' on payment of a fee of rupees fifty only.

7. holder of licence to keep account of quantities of opium. — Every person holding a licence in 'Form — O.P. II-A' shall keep in 'Form — G' daily accounts of the quantities of the medicines containing opium purchased & sold by him.

8. Licence for possessing opium for consumption. — (1) Any person desiring to possess opium for the purpose(s) of personal consumption shall make an application for a permit to the authorised officer.

(2) Subject to the provisions of sub-rule (3), on receipt of an application under sub-rule (1), the authorised officer, as the case may be, shall make such inquiries as he deems necessary and if he is satisfied that there is no objection to grant the permit applied for, he may subject to the orders of the Commissioner of Excise, if any, grant the applicant a permit in 'Form — O.P. III' on payment of a fee of rupees ten only.

Provided that no such permit shall be granted or renewed —
(i) except on the recommendation of the Medical Board or the Medical Officer, as the case may be, appointed in that behalf;
(ii) to a person under the age of twenty one years; or
(iii) to a person holding a permit in 'Form — B-III' appended to these rules.

(3) The authorised officer shall direct every applicant applying for a permit under this rule on grounds of addiction, to undergo medical treatment at the nearest hospital having facilities for the curative treatment of opium addicts in order to get rid of the addiction. He may, on the advice of the Medical Officer-in-charge of such Hospital grant or renew the permit to such applicant for the period recommended by such Medical Officer. Where the applicant fails to comply with the direction, the authorised officer may refuse to grant or renew the permit, unless —
(a) the applicant is over 60 years of age and his health does not permit undergoing hospitalisation; or
(b) the addiction of such person is certified by the Medical Board to be incurable even after undergoing the curative treatment at the hospital; or

(c) such person is the only earning member in his family and his hospitalization will, in the opinion of the authorised officer cause hardship to other members of his family so far as their maintenance is concerned.

9. Quantity of opium to be determined by authorised officer. — (1) A licence in 'Form — O.P.I.' or 'Form — O.P. II' shall be granted in respect of such quantity of opium as may be determined by the authorised officer in this behalf.

(2) A permit in 'Form — O.P. III' shall be granted in respect of such quantity of opium as may be fixed by the authorised officer, as the case may be, in accordance with the orders and directions issued by the Commissioner of Excise, from time to time;

Provided that the aggregate quantity that can be bought in a month shall not exceed 50 grams of opium and the quantity that can be possessed at any one time shall not exceed 5 grams of opium:

Provided further that the aggregate quantity that can be purchased in a month shall at every quarter of an year, except for special and adequate reasons, be subject to a reduction, the maximum being 12½ percent, and the minimum being ½ gram as the authorised officer may determine.

10. Licence not to be granted for any period beyond certain date. — No licence or permit under the aforesaid provisions shall be granted for any period beyond 31st March next following the date of the commencement of the licence/permit, as the case may be;

Provided that a licence in 'Form — O.P.I.' may be granted or renewed for a period not exceeding three years at a time but not beyond 31st March of the third year from the date of the commencement of the licence, if the licensee gives an undertaking to the effect that he shall abide by the provisions of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act No. 6 of 1985) and the rules, regulations and orders made thereunder and also by all the conditions of the licence which may be imposed by the Government at the time of granting the licence or at any time during the currency of the licence.

11. Depots for sale of opium. — Depots for the sale of opium shall be established at such places as the Government may from time to time direct.

12. Opium for sale at depot to be obtained from Uttar Pradesh. — Opium required for sale at a depot may be obtained from Gazipur in the Uttar Pradesh. Opium may also be obtained from such other places as the Government may direct.

13. Opium to be sold only at depot. — Opium shall not be sold at any place except at a depot established under rule 11:

Provided that a person holding a licence in 'Form — O.P.I.' or 'Form — O.P.II' or 'Form O.P. II-A' may sell medicines containing opium to the extent and subject to the conditions laid down in his licence.

14. Opium not to be imported by railway or by insured Postal Parcel. — Subject to the provisions of rule 12, no opium for the purposes of sale at a depot established under rule 11 shall be imported by railway or by Insured Postal Parcel;

Provided that a person holding licence in 'Form — O.P.I.' or 'Form O.P.II' or 'Form O.P.II-A' may, under a pass granted under sub-rule (2) of rule 15, import inter-state any medicines containing opium from any other State in India.

15. Interstate import of medicine, containing opium. — (1) Any person holding a licence in 'Form — O.P.I.' or 'Form O.P.II' or 'Form — O.P. II-A' desiring to import inter-state medicines containing opium from any other State in India shall make an application to the authorised officer in that behalf.

(2) On receipt of an application under sub-rule (1), the authorised officer shall make such inquiries as he deems necessary and if he is satisfied that there is no objection to grant the pass applied for, he may grant the applicant a pass in 'Form — A'.

16. Interstate export of medicine containing opium. — (1) Any person desiring to export inter-State any medicine containing opium shall make an application for a pass to the authorised officer in that behalf, and shall alongwith the application forward an import pass or a no-objection certificate from the Commissioner of Excise or the Chief Excise Authority of the place as the case may be, to which such medicines are to be exported.

(2) On receipt of an application under sub-rule (1), the authorised officer shall make such inquiries as he deems necessary and if he is satisfied that there is no objection to grant the pass applied for, he may, grant the applicant a pass in 'Form — B'.

(3) The authorised officer shall cause each package of medicines containing opium which is to be exported to be sealed with his official seal.

17. Application for pass to transport opium. — (1) Any person desiring to transport opium or any medicines containing opium shall make an application for a pass in that behalf as provided in sub-rule (2):

Provided that no such pass shall be necessary where transport of opium or any medicine containing opium is permitted under the licence or permit granted under these rules.

(2) Save as provided in sub-rule (1), a person holding a licence in 'Form — O.P.I.' or 'Form — O.P. II' or 'Form O.P. II-A' shall make an application to the authorised officer who granted him such licence. A person holding a permit in 'Form — O.P. III' shall make such application to the authorised officer who granted him such permit, and any other person shall make an application to the authorised officer of the place from which opium or any medicine containing opium is to be transported and shall in the case of a transport from one District to another
District, forward a no objection certificate of the authorised officer of such other District to which opium or any medicine containing opium is to be transported.

(3) On receipt of an application under sub-rule (2), the authorised officer, as the case may be, shall make such inquiries as he deems fit and if he is satisfied that there is no objection to grant the pass applied for, he may grant the applicant a pass in ‘Form—C’.

18. Packet containing opium not to be opened in transit.—No packet containing opium or any medicine containing opium shall be opened or broken while in transit:

Provided that nothing contained in this rule shall apply to the persons holding a permit in ‘Form—O.P.—III’:

19. Opium and pass for its transport to be examined by authorised officer at destination. — (1) Where opium or any medicine containing opium is transported from one District to another, the transport not being by railway administration shall, on arrival of the packages of opium or any medicine containing opium at their destination, present them together with the pass covering such transport for examination to the authorised officer.

(2) If after examination of the packages under sub-rule (1), the authorised officer is satisfied that the packages have not been tampered with in transit, and that the number of packages and their weight correspond with the number of packages and their weight specified in the pass, the packages together with the pass will be handed over to the transporter. If the authorised officer is not so satisfied, the packages shall be detained and dealt with according to the orders of the Commissioner of Excise.

MISCELLANEOUS

20. No railway administration shall.—

(a) receive or convey opium or any medicine containing opium which is not covered and accompanied by transport pass from an Officer duly empowered in this behalf; or

(b) convey opium or medicine containing opium otherwise than—

(i) in custody of a railway official up to the station at which the opium or any medicine containing opium should leave the railway, and

(ii) according to the route prescribed in such pass.

21. Bulk of consignment not to be broken in transit. — (1) The bulk of consignment of opium or any medicine containing opium in transit in the course of importation or exportation shall not be broken or opened any Revenue Officer, not below the rank of Manildar, or an Officer of the Drugs Controller’s Office, not below the rank of Inspector of Police may at any time examine such consignment.

(2) If, after such examination, the Officer is satisfied that the packages have not been broken, opened or tampered with in transit, and that the number of packages and their weight correspond with the number of packages and their weight specified in the pass, the consignment shall be allowed to proceed. If the said Officer is not so satisfied, the consignment shall be detained and dealt with according to the written orders of the Commissioner of Excise.

22. Preparation and admixtures to exhibit quantity of opium contained therein. — In the case of the preparations of admixtures containing opium (which are not manufactured drugs under the Act), the bottles, phials, packages or other containers of such preparations and admixtures or the labels affixed to them shall plainly exhibit,—

(a) the actual quantity of opium present in each such bottle, phial, package or container, or

(b) sufficient particulars thereof so as to admit the ready calculation of such quantity.

23. Any person may buy medicine containing opium prescribed by Registered Medical Practitioner. — Notwithstanding anything contained in these rules, it shall be lawful for any person to buy, possess, transport, consume or use any medicine containing opium in such quantity as may at one time be dispensed or sold to him in accordance with the prescription of a Registered Medical Practitioner.

24. Purchase of medicine containing opium on behalf of certain persons. — Any person may, without any permit or licence purchase, possess and transport opium or any medicine containing opium on behalf of a pardanishin lady, or on behalf of an infirm or invalid person who is physically not fit to purchase, possess and transport opium or any medicine containing opium provided that—

(i) the pardanishin lady or the infirm or invalid person holds a permit in Form O. P.—III or a prescription from a Registered Medical Practitioner for obtaining any medicine containing opium;

(ii) the person purchasing, possessing and transporting opium on behalf of the pardanishin lady or the infirm or invalid person otherwise than on a prescription has got a written authority in ‘Form-D’ from the pardanishin lady or the infirm or invalid person to do so on her or his behalf; and

(iii) the authorised officer empowered to grant permits in Form O.P-III has given his previous approval to such authority.

25. Purchase, etc. of opium on behalf of Government depots. — Notwithstanding anything contained in the foregoing rules, no licence, permit or pass shall be necessary for the purchase, possession, transport, import, export and sale of opium by or on behalf of Government depots established under rule 11.

26. Maintenance of monthly accounts of quantities of opium. — (1) Every person holding a licence in ‘Form-O.P.-I’, shall maintain in ‘Form-E’ monthly accounts of the quantities of opium purchased and used and of the balance held in stock by him.
(2) Every person holding a licence in 'Form-O.P.-II' shall maintain in 'Form-I' daily accounts of quantities of opium purchased and used by him.

CHAPTER III
Ganja

27. Prohibition of possession, etc of Ganja. — Possession, consumption and use, sale, export inter-state, import inter-state, transport of ganja and cultivation of any cannabis plant is prohibited.

CHAPTER IV
Medicinal opium manufacture

28. Prohibition of manufacture of medicinal opium. — The manufacture of medicinal opium is prohibited save under and in accordance with the conditions of a licence in 'Form-H' annexed hereto granted by the Drugs Controller Government of Goa, Daman and Diu.

CHAPTER V
Manufactured drugs

29. Drugs to be manufactured in accordance with conditions of licence. — No licensed dealer in manufactured drugs shall except in accordance with the conditions of his licence and except on the premises licensed for the purpose under these rules, manufacture any preparation containing any manufactured drugs from the materials which he is lawfully entitled to possess.

30. Manufactured drugs to be dispensed on prescriptions. — No licensed chemist shall dispense manufactured drugs except on prescription and in accordance with the conditions of his licence.

31. Possession of manufactured drug. — No person shall possess any manufactured drug except in such quantity as has been at one time dispensed or sold for his use in accordance with the provisions of rule 52 of these rules or of any corresponding rules for the time being in force in any part of India, the import inter-State or export inter-State of manufactured drugs save as has been permitted under the rules.

32. Possession of manufactured drugs by Registered Medical Practitioner. — (1) No Registered (Medical) practitioner shall for the purpose of sale, possess any quantity of manufactured drugs or any preparation containing any manufactured drugs:

Provided that such practitioner may, for use in his practice, possess —
(a) Opium derivatives containing in the aggregate not more than 4 grams of either morphine or diacetyl morphine or both;
(b) medicinal hemp not exceeding 30 grams of extract or 110 grams of tincture or both; and
(c) 1-methyl-4-Phenyl-piperidine-4-carboxylic acid ethyl ester (in the form of the hydrochloride, known under the names of Dolantin Demoral, Pethidine, Isopipacaine, etc) and its salts not exceeding 4 grams:

Provided further that the Licensing Authority may, by special order, authorise any such practitioner to possess as aforesaid any larger quantity of the said drugs.

(2) No registered (Medical) practitioner shall, for the purpose of sale, possess any quantity of coca derivatives:

Provided that such practitioner may under a special permit granted in this behalf by the Licensing Authority in "Form "NDPS" — I" hereto annexed, possess for use in his practice coca derivatives containing not more than 2 grams of cocaine in the aggregate:

Provided further that the Licensing Authority may authorise any such practitioner to possess as aforesaid a larger quantity of coca derivatives containing not more than 2 grams of cocaine.

(3) The expression "use in his practice" in sub rules (1) and (2) means only the actual direct administration of the drugs in injections, surgical operations or other emergent cases by or in the presence of Registered (Medical) practitioners. All other drugs issued by an approved practitioner, shall be deemed to be sales, except in the case of drugs issued free of charge from specially recognised charitable medical institutions.

33. Possession of manufactured drugs by Government Medical Officer. — (1) A Government Medical Officer in charge of Government and Government grant-in-aid Medical Institutions/hospitals/dispensaries/centres may possess manufactured drugs for use in such institutions.

(2) A registered (Medical) practitioner in charge of the Municipal Dispensaries or Incharge of hospitals and dispensaries belonging to missions and other corporate bodies and private nursing homes may possess manufactured drugs required for use in such dispensaries and hospitals.

(3) A Government Medical Officer-in-Charge of hospitals and dispensaries belonging to railways may possess manufactured drugs for use in such hospitals and dispensaries.

34. Medical Officer or Registered Medical Practitioner to keep accounts of manufactured drugs held in stock by him. — A Medical Officer or a Registered (Medical) practitioner possessing manufactured drugs under rule 33, shall:

(1) Keep accounts of manufactured drugs received, used and held in stock by him from time to time in the form prescribed by the Licensing Authority. The accounts shall be plainly and correctly written up daily in books bound, pagged and sealed with the seal of the Licensing Authority and shall show in each case of purchase, the date of purchase and the name and address of the person or firm from whom the purchase was made;

(2) Preserve the said accounts for not less than two years from the date of the last entry in the account book and shall produce them, together with any manufactured drugs that may be in his possession at the relevant time, for inspection on demand by the Licensing Authority, or by other officer duly authorised by him in this behalf;

(3) Furnish to the Licensing Authority or any other officer duly authorised by him in this
be held by him on the last day of the quarter in the form prescribed by the Licensing Authority, for the purpose.

35. Possession of opium derivatives and medical hemp. — (1) No person, unless he is authorised in this behalf by the Licensing Authority by an order made under sub-rule (1) of rule 55 shall possess opium derivatives and medical hemp not exceeding such quantities and otherwise than in such manner as may be specified in such order.

(2) No person, unless he is authorised in this behalf by the Licensing Authority by an order made under sub-rule (2) and (3) of rule 55 shall possess and use manufactured drugs for educational or scientific purposes and for use in an emergency and not exceeding such quantity and otherwise than in such manner as may be specified in such order.

36. Possession of manufactured drugs. — No licensed dealer in manufactured drugs or licensed chemists shall possess manufactured drugs except in such quantity and in such manner as may be specified in his licence.

37. Authorisation of inter-state import and export of manufactured drugs. — No person shall, without an authorisation granted to him under these rules for the import inter-state; export inter-state or transport of manufactured drugs, possess such drugs exceeding such quantity and otherwise than in such manner as may be specified in such authorisation.

38. Import Inter-State, Export Inter-State and Transport. — No person shall import inter-state, export inter-state or transport manufactured drugs except in such quantities as he may lawfully possess under rule 31.

39. (1) Registered (Medical) practitioner shall import inter-state, export inter-state or transport opium derivatives, medicinal hemp and 1-methyl-3-phenyl-piperidine-4-carboxylic acid ethyl ester (in the form of the hydrochloride known under the names of Donlatin, Demerol, Phethidine, Isoniazide, etc.) and its salts except in such quantities as he may lawfully possess under sub-rule (1) of rule 32.

(2) No practitioner shall, without a licence in 'Form NDPS — 2' hereto annexed or a special permit in 'Form NDPS — 3' hereto annexed, import inter-state, export inter-state or transport coca derivatives containing not more than 1,300 milligrams of cocaine for use in his practice, provided that no such export inter-state or transport shall be made for sale.

40. Transport of manufactured drugs by Medical Officer. — (1) A Medical Officer-in-charge of hospitals and dispensaries belonging to the railways may transport manufactured drugs required for use in such hospitals and dispensaries.

(2) A Medical Officer-in-charge of Government or Government grant-in-aid Medical Institution may transport manufactured drugs required for use in such institutions.

41. Inter-state import and transport of opium derivatives. — (1) No person unless he is authorised in this behalf by the Licensing Authority by an order made under sub-rule (1) of rule 55 shall import inter-state or transport opium derivatives or medicinal hemp and not exceeding such quantities and otherwise than in such manner as may be specified in such order.

(2) No person, unless he is authorised in this behalf by the Licensing Authority by an order made under sub-rule (2) of rule 55 shall transport manufactured drugs for educational or scientific purposes and not exceeding such quantities and otherwise than in such manner as may be specified in such order.

42. Inter-state import not to exceed authorised quantity. — No person shall, without an authorisation granted to him under these rules for the import inter-state of any manufactured drug import that drug exceeding such quantity and otherwise than in such manner as may be specified in such authorisation.

43. Conditions for import of medicinal opium. — No licensed dealer in manufactured drugs shall, unless permitted by the Licensing Authority import inter-state medicinal opium from the Ghazipur Factory Uttar Pradesh (hereinafter referred to as "the factory"). Such import inter-state shall be permitted subject to the following conditions:

(1) Every application by the Licensed dealer for the supply of medicinal opium shall be sent to the Licensing Authority.

(2) The stock in hand on the date of the application shall be stated and the quantity of opium applied for shall not be more than sufficient for six months manufacture, subject however, to the minimum of 1 kilogram prescribed in condition (9) for individual indents.

(3) The purposes for which the medicinal opium is required shall be specified in the application.

(4) An intimation of the despatch of each consignment shall be sent by the Superintendent of the Factory to the Licensing Authority who shall make arrangements for its examination on arrival by a responsible officer not below the rank of Drug Inspector of the Office of Drugs Controller.

(5) A complete record of the quantity of medicinal opium received and used for each preparation, with the amount of each product manufactured and its morphine contents shall be maintained in such manner as the Licensing Authority may prescribe "such record shall be periodically examined by any Officer not below the rank of Drugs Inspector".

(6) Every consignment on arrival shall be examined by the officer referred to in condition (4) and immediately brought to account in the form prescribed in condition (5). No part of the
medicinal opium shall be medicated except in the presence of such officer, who shall witness—

(a) in the case of liquid extracts or tinctures the beginning of the panning in the evaporators or the addition of the solvent in the percolators;

(b) in the case of mixtures the thorough admixture of the medicinal opium with other drugs.

Twelve clear days notice shall be given to such officer of any medication of medicinal opium.

(7) No medicinal opium shall under any circumstances be sold or shall be allowed to be removed from the premises of the person to whom it has been issued, otherwise than as a part of a manufactured medicinal preparation, and the medicinal opium shall be used for the purpose other than those specified in the application.

(8) It shall be competent for the Government to supply medicinal opium to such persons as it deems fit.

(9) The limit of the amount of medicinal opium to be supplied to any person in any calendar year shall be 10 kilograms and individual indent shall not be for less than 1 kilogram or more than 5 kilograms at a time.

44. Inter state export of manufactured drugs.—No licensed dealer in manufactured drugs shall, except under an authorization granted under rule 59 and subject to the conditions of his licence, export inter-state manufactured drugs to any part in India.

45. Inter state export of opium derivatives.—No person, unless he is authorised in this behalf by the Drugs Controller's Office by special order made under “rule 56” shall export inter-state opium derivatives or medicinal hemp exceeding such quantity and otherwise than in such manner as may be specified in such order.

46. Transport of manufactured drugs not to exceed authorised quantity.—No person shall, without an authorisation granted to him under these rules for the transport of manufactured drugs, transport such drug exceeding such quantity and otherwise than in such manner as may be specified in such authorisation:

Provided that Registered (Medical) practitioners holding a licence in 'Form NDPS-2' may, without such authorisation, transport otherwise than by rail or post manufactured drugs in quantities not exceeding those specified in condition 1 of his licence.

47. Inter state import or export to be subject to directions of Drugs Controller.—Every person importing inter-state, exporting inter-state or transporting manufactured drugs shall comply with such general or special directions as may, from time to time, be issued by the Drugs Controller.

48. Inter state import or export of manufactured drugs in Goa, Daman and Diu.—Except as provided in rule 49, no person shall import inter-state or export inter-state or transport manufactured drugs whether or not the same is in the possession of Government by rail or post into, out of or within the Union territory.

49. Conditions for inter state import or export of manufactured drugs.—Import inter-state, export inter-state or transport of manufactured drugs shall be allowed by rail or post subject to the following conditions, namely:

(a) the parcel of the manufactured drugs when sent by post shall be sent by registered parcel only;

(b) the parcel of such drugs whether sent by rail or by post shall be insured;

(c) the parcel shall be covered by a permit issued in this behalf by the competent authority at the place to which the parcel is addressed;

(d) the parcel shall be accompanied by a declaration showing the names of the consignor and the consignee, the contents of the parcel in detail, the number and date of the permit covering the import inter-state, export inter-state, or transport, as the case may be, and the number of the licence, if any, held by the consignor or the consignee;

(e) the consignor and the consignee, if they are licencsees, shall show distinctly in their account books, the names of the consignee and the consignor, respectively, and the quantities of the drugs imported inter-state, exported inter-state or transported by and to them, as the case may be from time to time by post.

50. Miscellaneous.—Nothing in these rules shall be deemed to permit the import inter-state of manufactured drugs from any parts of India outside the Union territory unless the rules for the time being in force in such parts of India relating to the export inter-state have been complied with.

51. Conditions for sale of manufactured drugs.—

(1) A Licensed Dealer in manufactured drugs may sell, otherwise than on prescription manufactured drugs subject to the conditions of his licence.

(2) Such dealer shall maintain a written record of every sale made under the licence in the manner laid down therein and in such manner as the Drugs Controller's Office may from time to time direct, and shall preserve such record for not less than two years from the date of the last entry therein.

52. Licensed chemist to sell manufactured drugs on prescription.—No licensed chemist shall sell manufactured drugs otherwise than on prescription and subject to the condition of his licence.

53. Sale of manufactured drugs to Government office.—Notwithstanding anything contained in these rules, holder of a licence in 'Form NDPS-1' shall, whenever required to do so, sell any manufactured drugs to any Government Officer who is duly authorised by the Government in this behalf to possess such drug on behalf of Government under the provisions of section 10 of the Act, provided that a receipt is obtained by the holder of the licence from such officer for the same and kept on record.
54. Conditions for prescription of manufactured drugs.—A prescription for the supply of manufactured drugs shall comply with the following conditions:

(1) The prescription shall be in writing dated and signed by a Registered (Medical) practitioner with his full name, qualifications and correct address. The prescription shall also specify the name and correct address of the person to whom such prescription is given and the total quantity of drugs to be supplied thereon.

(2) The prescription shall not be given for the use of the prescriber himself or any member of his family;

(3) A prescription given by a registered Dentist shall be only for the purpose of dental treatment and shall be marked “For Local Dental Treatment Only”.

(4) A prescription given by an approved Veterinary Surgeon shall be only for the purpose of treatment of animals and shall be marked “For Animal Treatment Only”.

(5) No practitioner shall give any prescription for the supply of any of the manufactured drugs otherwise than in accordance with the foregoing conditions.

55. Authorisation by Drugs Controller.—The Drugs Controller may by a general or special order, authorise

(1) any person in-charge of an educational institution or engaged in scientific research to possess and use, for educational and/or scientific purposes only, manufactured drugs in such quantity and in such manner as may be specified in the said order.

(2) The Pilot of an Aircraft to possess and use on the Aircraft in an emergency preparations containing morphine in such quantity and in such manner as may be specified in such order;

(3) A person in charge of an ambulance or a first aid station or a first-aid box to possess and use in an emergency the manufactured drugs in such quantity and in such manner as may be specified in such order.

56. Authorisation by Drugs Controller for inter state opium derivatives.—The Drugs Controller may by a special order authorise any person to export inter-state opium derivatives or medicinal hemp, subject to such conditions as may be specified in such order.

57. Granting of dealer’s licence.—(1) The Licensing Authority or any other officer empowered in this behalf by the Drugs Controller may grant a dealer’s licence in “Form NDPS-1” or a chemist’s licence in “Form NDPS-2”, to any person who in the opinion of the Licensing Authority or such officer, as the case may be, is not likely to abuse such grant, and may determine the quantity of the drugs to be possessed under such licences.

(2) The fees as shown below shall be charged in advance to the holders of a licence in ‘Form NDPS-1’ and Form NDPS-2:

<table>
<thead>
<tr>
<th>Form</th>
<th>Fee</th>
</tr>
</thead>
<tbody>
<tr>
<td>N.D.P.S. 1</td>
<td>Rupees 20 per annum</td>
</tr>
<tr>
<td>N.D.P.S. 2</td>
<td>Rupees 20 per annum</td>
</tr>
</tbody>
</table>

Provided that for the purpose of charging the fees the fraction of a year shall be reckoned as one complete year.

(3) No licence under sub-rule (1) shall be granted for a period extending beyond 31st December next following the date of the commencement of the licence;

Provided that a licence in Form — N.D.P.S. 2 may be granted or renewed to a Registered Medical practitioner for a period not exceeding three years at a time but in no case shall the period end beyond 31st December of the third year from the date of the commencement of licence;

Provided further that the holder of a licence shall have to abide by the provisions or the Act, regulations, rules and orders made thereunder and the conditions of the licence and also by such other conditions of the licence as may be prescribed by the Government during the currency of the licence and shall give an undertaking to that effect.

58. Authorisation for inter state import of manufactured drugs.—The Licensing Authority or such officer, as the Licensing Authority may empower in this behalf, may grant to any licensed dealer in manufactured drugs, or a licensed chemist an authorisation in ‘Form-NDPS.4’ for the import inter-state of manufactured drugs not exceeding the quantity which such dealer or chemist may lawfully possess.

59. No objection certificate.—When any manufactured drugs are to be exported to any other part of India, the person intending to export the same shall first obtain a no-objection certificate or permit from an officer authorised in this behalf under the relevant rules in force in such other part of India and present such certificate or permit along with the indent, to the Licensing Authority or such officer as may be authorised in this behalf, at the place of export, who shall then, if he sees no objection, issue an export authorisation in ‘Form -NDPS.5’;

Provided that in the case of export inter-state of the drugs required for use in Government Medical Institutions no such no-objection certificate or permit shall be necessary if the Director of Health Services or the Director of the Government Veterinary Institute in the District to which the drug is to be exported, certifies by endorsement on the requisition or indent that the drug is required for bona fide Government purpose and intimates the facts to the Drugs Controller and the concerned authority of the place from and to which the drugs are to be exported.

60. Transport authorisation.—When any manufactured drug is to be transported, the person intending to transport the same shall first obtain a transport authorization in ‘Form NDPS.6’ from the Licensing Authority or such other officer as may be authorised by the Drugs Controller in this behalf,
at the place to which the drug is to be transported and present it to the Licensing Authority or such other Officer as may be authorised by the Drugs Controller in this behalf, at the place from which the drug is to be transported, who shall complete the authorization and allow the removal of the drug, provided that the quantity of the drug does not exceed the quantity which such person may lawfully possess.

61. Special authorisation.—(1) The Licensing Authority may grant a special authorisation in Form—NDPS.3 hereto annexed to a registered (Medical) practitioner for the possession and use in the exercise of his practice, but not for sale of cocoa derivatives containing not more than one gram of cocaine;

Provided that the Licensing Authority may allow a larger quantity of the drug containing not more than one gramme of cocaine in such cases as he may, having regard to the requirements of the permit holder, consider fit.

(2) The special authorization in Form—NDPS.3 may be granted or renewed for a period not exceeding three years at a time but in no case shall extend beyond the 31st December of the third year from the date of commencement of the authorization.

62. Limits to be specified with reference to manufactured drugs.—In the case of preparation and admixtures containing manufactured drugs, the limit wherever specified shall be with reference to the manufactured drug contents, respectively, and not with reference to the quantity or bulk of the preparation and the bottles, phials, packages or other containers of these preparations and labels affixed to them shall plainly exhibit the actual quantity of the manufactured drugs present in such container or sufficient particulars to admit for the ready calculation of such quantity.

63. Certain preparations not to be declared manufactured drug.—All preparations containing not more than 0.2 percent of morphine or 0.1 percent of cocaine and any other preparations which the Central Government may by notification in the Gazette of India make in pursuance of a finding under Article 8 of the Geneva Convention declare not to be manufactured drug, may be imported, exported, transported, possessed or sold without any restriction.

64. Directions by Government.—Notwithstanding anything contained in these rules the Government may, by notification in the Official Gazette, direct that no licence, permit, pass or authorization shall be necessary for the import, export, transport, possession or sale, in quantity not exceeding 500 grams of such manufactured drugs as may be specified in the notification.

CHAPTER VI
Rewards

65. Rewards to officials.—(1) In cases involving the seizure of opium and other narcotic drugs, reward upto 10% of the estimated market value of the goods involved (half of the maximum reward) indicated in the annexure as amended from time to time, in respect of opium and other narcotic drugs and Psychotropic substances alongside fine imposed may be granted in such proportion as the Commissioner of Excise/Inspector General of Police/Drugs Controller may think fit to any official of the Excise/Police/Office of Drugs Controller respectively;

Provided rewards in excess of the above limit, not exceeding 20% (as shown in the annexure and as revised from time to time) of the said value of illicit prices indicated, alongwith fine imposed may be considered in cases where the Government servant has exposed himself to great personal hazard or displayed exemplary courage, initiative, ingenuity or resourcefulness of an extra-ordinary character of his personal efforts have been mainly responsible for the detection of the contraband, narcotic drugs/ Psychotropic Substances. The prescribed purity as indicated in the Annexure will be determined after getting the potency of the seized drug tested at the Combined Food and Drugs Laboratory of the Directorate of Health Services for the payment of reward in question. However, 50% of the total amount of reward admissible may be disbursed as soon as it is identified that the seized drug is a narcotic drug.

Provided further that rewards exceeding Rs. 50,000/- but upto Rs. 1,00,000/- shall be granted by the concerned Secretary to Government and over Rs. 1,00,000/- shall be decided and granted by a committee consisting of—

(i) The Chief Secretary.
(ii) The Secretary of the Department.
(iii) The Departmental Head.

(2) Half of the reward shall be given to the informer, if any, and the remainder to be distributed to the persons who actually co-operated in the seizure or arrest.

(3) If there is no informer, the whole reward shall go to the persons who actually co-operated in the seizure or arrest.

(4) Advance of rewards will be paid to the informers and Government servants upto 50% of the expected final reward immediately after the seizure.

(5) In exceptional cases; the Head of the Department may, having regard to the value of the seizures effected and magnitude of the crime detected and special efforts or ingenuity displayed by the officers concerned, sanction and announce the grant of suitable rewards on the spot to be adjusted against the advance reward that may be sanctioned.

(6) Final reward will be paid after adjusting the advance reward, if any, paid only after the case is finalised by appropriate courts of law leading to confiscation of the contraband and/or conviction of the accused and/or imposition of fines.

(7) No reward shall ordinarily be granted for officer above the rank of Group 'A' Officer equivalent to Commissioner of Excise. However, in appropriate Cases the Government may consider the grant of lumpsum payment/advance increments and/or recognition in any other manner of the service rendered, to officers above Group 'B'.

(8) The rewards shall be equally distributed to the eligible Government servants in proportion to the risk and responsibility undertaken by them.
(9) The Commissioner/Drugs Controller may incur an expenditure not exceeding Rs. 250/- in each case for the employment of informer or for any other purpose connected with the prevention or detection of any offence under the Act.

(4) The fee payable for each amendment to be made in a licence, authorisation, permit or pass shall be rupees five only.

67. Authority granting licence may cancel the same.— (1) The authority granting any licence, permit, pass or authorization under these rules may, for the reasons to be recorded in writing, cancel or suspend the same,

(a) if the purpose for which the licence, permit, pass or authorization was granted ceases to exist;

(b) in the event of any breach by the holder or by his servant or by any person acting with his express or implied permission on his behalf, of such licence, permit, pass or authorisation or any of the terms and conditions thereof or of any licence, permit, pass or authorisation as the case may be, previously held by him.

68. Authority granting licence may cancel it. Whenever the authority granting a licence, permit, pass or authorization, considers that it should be cancelled or suspended for any cause other than those specified in rule 67 may cancel or suspend it by recording the reasons in writing for doing so.

69. Appeals. — (1) All orders passed by an Authorised Officer under these rules shall be appealable to the Chief Secretary of Goa, Daman and Diu.

(2) All orders passed by the Licensing Authority shall be appealable to the Chief Secretary of Goa, Daman and Diu.

(3) Every appeal under these rules shall be filed within a period of thirty days from the date of the order appealed against and shall be accompanied by a certified copy of the said order.

70. Supplementary matters. — (i) The Commissioner of Excise may issue written instructions for providing any supplementary matters arising out of chapter II of these rules.

(ii) The Drugs Controller may issue written instructions providing for any supplementary matters arising out of chapter IV and V of these rules.

---

**CHAPTER VII**

**Miscellaneous**

66. Fees. — (1) The fee payable in respect of transfer of a licence from one place to another shall be the same as payable for grant of a licence.

(2) The fee payable in respect of the transfer of a licence from one name to another shall be the same as payable for grant of a licence.

(3) The fee payable for supply to a licensee of a duplicate copy of a licence, authorization, permit or pass shall be rupees five only.

---

**ANNEXURE**

<table>
<thead>
<tr>
<th>St. No.</th>
<th>Commodity</th>
<th>Prescribed purity</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Opiate</td>
<td>Standard opium, 90% or more of morphine.</td>
</tr>
<tr>
<td>2</td>
<td>Morphine base and its salts</td>
<td>90% or more of dihydrocodeine.</td>
</tr>
<tr>
<td>3</td>
<td>Heroin and its salts</td>
<td>90% or more of dihydrocodeine.</td>
</tr>
<tr>
<td>4</td>
<td>Hashish</td>
<td>90% or more of dihydrocodeine.</td>
</tr>
<tr>
<td>5</td>
<td>Hashish oil</td>
<td>With THC content of 20% or more.</td>
</tr>
<tr>
<td>6</td>
<td>Gauna</td>
<td>Should be commercially acceptable as per above.</td>
</tr>
</tbody>
</table>

---

**FORM O.P.I.**

[See rule 4(2)]

Licence No. ...

Licence for the possession of opium for use as an ingredient of any medicine and for the sale of medicines containing opium on prescription

Licence is hereby granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules, regulations and orders made thereunder to Shri/Smt./Kum./M/s ... of ... (hereafter called "the licensee") to purchase, transport and possess opium and to use it as an ingredient of any medicine and to sell the medicines containing opium in his dispensary situated at ... in the Taluka of ... in the ... District, subject to the following conditions, namely:

**CONDITIONS**

1. This licence shall remain in force from ... to ... (both days inclusive).

2. The licence shall not obtain opium except from a depot established under rule 4.1 of the Goa, Daman and Diu Narcotics Drugs and Psychotropic Substances Rules, 1987.
Provided that the licensee may, in any special case, be permitted to obtain opium from any other place with the previous approval of the Commissioner of Excise:

Provided further that the licensee may obtain his requirements of medicines containing opium from any licensee who is permitted to sell such medicines under Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987, or may import the same from any other State in India, subject to the provisions of the said rules.

3. (1) The licensee shall not purchase, during any quarter commencing from the 1st day of April, opium exceeding 3 grams and shall not possess at any time in excess of double the quantity:

Provided that where the licensee fails to purchase any quantity of opium which he is authorised to purchase during a quarter, he shall not be entitled to purchase the same at any time thereafter during the licence period except during the quarter next following.

(2) The licensee shall get the details of the purchase entered in Schedule hereto annexed by the Officer-in-charge of the depot before he removes from the depot the opium purchased by him.

4. The licensee shall not use or sell opium except as an ingredient of any medicine prescribed and dispensed by him for his patients. He shall not dispense any medicine containing opium except under a prescription issued by him and in the manner laid down in such prescription.

5. The licensee shall not keep opium and any medicines containing opium except at his dispensary.

6. No opium other than the opium obtained under this licence shall be transported, possessed or used by the licensee.

7. The privileges of purchase, possession and transport of opium granted under this licence shall extend only so far as they are incidental to its use in accordance with this licence.

8. (1) The licensee shall keep monthly accounts of the quantities of opium purchased and used and of the balance held in stock by him in the form prescribed by rule 26 of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987. The accounts shall be plainly and correctly written up in a bound book, paged and stamped with the seal of the Superintendent of Excise, similar accounts in respect of any medicine containing opium prepared, purchased or imported by the licensee shall also be maintained from day to day in such form as may be prescribed by the Government.

(2) The licensee shall file and preserve for two years the said accounts, passes and the prescriptions, in original in which he has prescribed opium as one of the ingredients of any medicines and shall produce them for inspection, along with the opium held by him in balance, at any time when the Authorised Officer calls upon him to do so.

9. The licence may be suspended or cancelled in accordance with the provisions of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987.

10. In case this licence is cancelled during the currency of the period for which it is granted or is not renewed on its expiry, the licensee shall forthwith hand over the whole of the unused stock of opium and of any medicines containing opium to the authorised officer.

The licensee shall also hand over to the authorised officer all accounts, passes and prescription in original, which he is required to keep and preserve under this licence.

Granted this ... day of ... 19 ... 

- Seal -

Authorised Officer

Place: 

SCHEDULE

Licence No. ...

Name of the depot: 

Name of the licensee: 

Full Address: 

---

Aggregate quantity of opium allowed to be purchased during each quarter

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Date</th>
<th>Quantity purchased</th>
<th>Progressive total of purchases for each quarter</th>
<th>Signature with date of the officer-in-charge of the depot</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>

FORM - O. P. II

(See rule 5 (2))

Licence No. ...

Licence for the possession of opium required on behalf of any institution, or by the manufacturer of medicines, for use as an ingredient of any medicines and for the sale of medicines containing opium

Licence is hereby granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules, regulations, and orders made thereunder to Shri/Smt./Kum/M/s ..., (hereinafter called "the licensee") in respect of (hereinafter called "the said institution/manufacturer/Chemist"), on payment of a licence fee of Rs. ..., authorising him to buy, transport and possess opium and to use it as an ingredient of any medicines and to sell the medicines containing opium at the premises of the said institution/manufacturer/Chemist situated at ..., in the ..., District subject to the following conditions, namely:

CONDITIONS

1. This licence shall remain in force from ... to ... (both days inclusive).

2. The licensee shall not obtain opium except from a depot established under rule 11 of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987 or may import the same from any other place with the previous approval of the Commissioner of Excise:

Provided that the licensee may, in any special case, be permitted to obtain opium from any other place with the previous approval of the Commissioner of Excise:

Provided further that the licensee may obtain his requirement of any medicines containing opium from any licensee who is permitted to sell such medicines under Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987 or may import the same from any other State in India, subject to the provisions of the said Rules.

3. (1) The licensee shall not purchase, during any quarter commencing from the 1st day of April, opium exceeding 3 grams and shall not possess it at any time in excess of double the quantity.

Provided that where the licensee fails to purchase any quantity of opium which he is authorised to purchase during a quarter, he shall not be entitled to purchase the same at any time thereafter during the licence period except during the quarter next following.

(2) The licensee shall get the details of the purchase entered in Schedule hereto appended by the Officer-in-charge of the depot, before he removes from the depot the opium purchased by him.

4. The licensee shall not use or sell opium except as an ingredient of any medicine prescribed and dispensed by him for his patients. He shall not dispense any medicine containing opium except under a prescription issued by him and in the manner laid down in such prescription.

5. The licensee shall not keep opium and any medicines containing opium except at his dispensary.

6. No opium other than the opium obtained under this licence shall be transported, possessed or used by the licensee.

7. The privileges of purchase, possession and transport of opium granted under this licence shall extend only so far as they are incidental to its use in accordance with this licence.

8. (1) The licensee shall keep monthly accounts of the quantities of opium purchased and used and of the balance held in stock by him in the form prescribed by rule 26 of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987. The accounts shall be plainly and correctly written up in a bound book, paged and stamped with the seal of the Superintendent of Excise, similar accounts in respect of any medicine containing opium prepared, purchased or imported by the licensee shall also be maintained from day to day in such form as may be prescribed by the Government.

(2) The licensee shall file and preserve for two years the said accounts, passes and the prescriptions, in original in which he has prescribed opium as one of the ingredients of any medicines and shall produce them for inspection, along with the opium held by him in balance, at any time when the Authorised Officer calls upon him to do so.

9. The licence may be suspended or cancelled in accordance with the provisions of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987.

10. In case this licence is cancelled during the currency of the period for which it is granted or is not renewed on its expiry, the licensee shall forthwith hand over the whole of the unused stock of opium and of any medicines containing opium to the authorised officer.

The licensee shall also hand over to the authorised officer all accounts, passes and prescription in original, which he is required to keep and preserve under this licence.

* Strike out the words which are not applicable.

---

5. The licensee shall not keep opium and medicines containing opium except at the premises of the said institution/manufacturer/Chemist.

6. No opium other than the opium obtained under this licence shall be transported, possessed or used by the licensee.
The privileges of purchase, possession and transport of opium granted under this licence shall extend only so far as they are incidental to its use in accordance with this licence.

8. (1) The licensee shall keep daily accounts of the quantities of opium purchased and used by him in the form prescribed by rule 26 of the Goa, Daman and Diu Narcotics Drugs and Psychotropic Substances Rules, 1987. The accounts shall be plainly and correctly written up in a bound book, pagd and stamped with the seal of the Superintendent of Excise. Similar accounts in respect of any medicines containing opium prepared, purchased or imported by the licensee shall also be maintained from day to day in such form as may be prescribed by the Government.

(2) The licensee shall file and preserve for two years the said accounts, passes and copies of the prescriptions against which he sold the opium as one of the ingredients of any medicines and shall produce them for inspection along with the opium and the medicines containing opium held by him in balance, at any time when the authorised officer calls upon him to do so.

9. Except with the permission of the authorised officer, the licensee shall not sell the privileges conferred upon him by this licence nor shall he admit any person as his partner in the business of his manufacture.

10. This licence may be suspended or cancelled in accordance with the provisions of the Goa, Daman and Diu Narcotics Drugs and Psychotropic Substances Rules, 1987.

11. In case this licence is cancelled during the currency of the period for which it is granted or is not renewed on its expiry, the licensee shall forthwith hand over the whole of the unused stock of opium and of the medicines containing opium to the authorised officer. The licensee shall also hand over to the authorised officer, all accounts passes and prescriptions in original which he is required to keep and preserve under this licence.

Granted this ... day of ... 19...

— Seal —  
Authorised Officer

Place:

SCHEDULE

Licence No.

Name of the licensee:

Full Address:

Aggregate quantity of opium allowed to be purchased during each quarter

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Date of purchase</th>
<th>Quantity purchased</th>
<th>Progressive total of purchases for each quarter</th>
<th>Signature with date of the officer</th>
<th>In-charge of the depot</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

FORM — O.P.II-A
(See rule 7(2))

Licence for the possession and sale of medicines containing opium by a dealer

Licence is hereby granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules, regulations and orders made thereunder to Shri/Smt./Kum./Ms. ... (hereinafter called "the licenise") on payment of licence fee of Rs. authorising him to buy, transport, possess and sell any medicines containing opium at his premises situated at ... in the Taluka of ... in the District subject to the following conditions namely:

CONDITIONS

1. This licence shall remain in force from ... to ... (both days inclusive).

2. The licensee shall not obtain any medicines containing opium except as permitted under the Goa, Daman and Diu Narcotics Drugs Psychotropic Substances Rules, 1987.

3. The licensee shall not sell any medicines containing opium to any person other than a person holding:

(i) a licence in Form O.P.I. or Form O.P.II, or Form O.P.II-A:

(ii) a prescription issued by a registered medical practitioner in that respect; nor shall he sell such medicines in a manner different from that laid down in such licence or prescription.

4. The licensee shall not keep any medicines containing opium except at the above said premises.

5. No medicines containing opium other than those obtained under this licence shall be transported, possessed or sold by the licensee.

6. The licensee shall keep daily accounts of the quantities or medicines containing opium purchased and sold by him in 'Form — O'. The accounts shall be plainly and correctly written up in a bound book, pagd and stamped with seal of the Superintendent of Excise.

7. The licensee shall file and preserve for two years the said accounts, passes and copies of the prescriptions against which he sold any medicines, containing opium and shall produce them for inspection along with the stock of the medicines containing opium held by him in balance at any time when the authorised officer or any other officer duly empowered in this behalf calls upon him to do so.

8. Except with the permission of the authorised officer the licensee shall not sell, transfer or sublet the privileges conferred upon him by this licence nor shall he admit any person as his partner in the business of his licence.

9. This licence may be suspended or cancelled in accordance with the provisions of Goa, Daman and Diu Narcotics Drugs and Psychotropic Substances Rules, 1987.

10. In case this licence is cancelled during the currency of the period for which it is granted or is not renewed on its expiry, the licensee shall forthwith hand over the whole of the unused stock of opium and of the medicines containing opium to the authorised officer. The licensee shall also hand over to the authorised officer, all accounts passes and prescriptions in original which he is required to keep and preserve under this licence.

Granted this ... day of ... 19...

— Seal —  
Authorised Officer

Place:

[Confidential]

FORM — O.P.III
(See rule 8(2))

Permit for the possession of opium for personal consumption in the Union territory of Goa, Daman and Diu

(A) Purpose for which the permit is granted.

(1) Name and address of the Medical Board which granted the certificate.

(B) Reference to medical certificate.

(1) Name and address of the Medical Board which granted the certificate.

(2) Date of certificate.

(3) Drug recommended.

(4) Quantity recommended per month.

(5) Personal identification marks of the permit holder as verified by the Medical Board.

(C) Reference to medical certificate.

(1) Name and address of the Medical Board which granted the certificate.

(2) Date of certificate.

(3) Drug recommended.

(4) Quantity recommended per month.

(5) Personal identification marks of the permit holder as verified by the Medical Board.

(place to be filled by the Superintendent of Excise in the State/Union Territory of Goa, Daman and Diu)
This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules, regulations and orders made thereunder to ... to ... (hereinafter referred to as “the permit holder”), on payment of a fee of Rs. ... authorising him to buy, possess, transport and consume opium subject to the following conditions:—

**CONDITIONS**

1. This permit shall remain in force from ... to ... (both days inclusive).
2. The permit holder shall as soon as possible present this permit before the Inspector of Excise, for his counter signature and in any case not later than one month from the receipt of this permit.
3. (1) The permit holder shall not purchase during any one month opium exceeding 5 grams:
   Provided that his quantity may be reduced during the currency of the permit in accordance with the provision in Goa, Daman and Diu Narcotics Drugs and Psychotropic Substances Rules, 1987.
   (2) The permit holder shall not possess at any one time more than 5 grams of opium.
4. (1) The permit holder shall not obtain his supplies of opium from any place except from a depot established under rule 11 of Goa, Daman and Diu Narcotics Drugs and Psychotropic Substances Rules, 1987.
   (2) The permit holder shall get the details of the purchase entered on the reverse of the permit by the Officer-in-charge of the depot, before he removes from the depot the opium purchased by him.
5. No opium other than opium obtained under this permit shall be transported possessed or consumed by the permit holder.
6. The privileges of purchase, transport and possession of opium granted under this permit shall extend only so far as they are incidental to its consumption in accordance with this permit.
7. The permit shall be non-transferable and may be suspended or cancelled in accordance with the provisions of the Goa, Daman and Diu Narcotics Drugs and Psychotropic Substances Rules, 1987.
8. In case the permit is cancelled during its currency or is not renewed on its expiry, the whole of the un consumed stock of opium shall forthwith be surrendered to the authorised officer.

**Form A**

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:—

1) The consignment shall not be opened or broken in transit.
2) This pass shall remain in force up to and including ... 19 ...

---

**Pass for import inter-State of medicine(s) containing opium**

<table>
<thead>
<tr>
<th>Serial No.:</th>
<th>Dated:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shri/Smt/Kum/M/s.</td>
<td>* ......... is hereby authorised to import inter-State the undermentioned medicines containing opium from Shri/Smt/Kum/M/s. + ...........</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be imported</th>
<th>Total quantity of opium contained in the medicines to be imported</th>
<th>Packages Gross No. weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>*</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

---

**Form A**

[See rule 15(2)]

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:—

1) The consignment shall not be opened or broken in transit.
2) This pass shall remain in force up to and including ... 19 ...

---

**Pass for Import inter-State of Medicine(s) containing opium**

<table>
<thead>
<tr>
<th>Serial No.:</th>
<th>Dated:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shri/Smt/Kum/M/s.</td>
<td>* ......... is hereby authorised to import inter-State the undermentioned medicines containing opium from Shri/Smt/Kum/M/s. + ...........</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be imported</th>
<th>Total quantity of opium contained in the medicines to be imported</th>
<th>Packages Gross No. weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>*</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

---

**Form A**

[See rule 15(2)]

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:—

1) The consignment shall not be opened or broken in transit.
2) This pass shall remain in force up to and including ... 19 ...

---

**Pass for Import inter-State of Medicine(s) containing opium**

<table>
<thead>
<tr>
<th>Serial No.:</th>
<th>Dated:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shri/Smt/Kum/M/s.</td>
<td>* ......... is hereby authorised to import inter-State the undermentioned medicines containing opium from Shri/Smt/Kum/M/s. + ...........</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be imported</th>
<th>Total quantity of opium contained in the medicines to be imported</th>
<th>Packages Gross No. weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>*</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

---

**Form A**

[See rule 15(2)]

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:—

1) The consignment shall not be opened or broken in transit.
2) This pass shall remain in force up to and including ... 19 ...

---

**Pass for Import inter-State of Medicine(s) containing opium**

<table>
<thead>
<tr>
<th>Serial No.:</th>
<th>Dated:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shri/Smt/Kum/M/s.</td>
<td>* ......... is hereby authorised to import inter-State the undermentioned medicines containing opium from Shri/Smt/Kum/M/s. + ...........</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be imported</th>
<th>Total quantity of opium contained in the medicines to be imported</th>
<th>Packages Gross No. weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>*</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>
FORM — A

[See rule 15(2)]

Pass for import inter-State of medicine(s) containing opium

(To be forwarded to the Excise Authority of the place of export).

Serial No.:

Dated:

Shri/Smt./Kum./M/s. * .......... is/are hereby authorised to import inter-State the undermentioned medicines containing opium from Shri/Smt./Kum./M/s. + .......... 

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be imported</th>
<th>Total quantity of opium contained in the medicines to be exported</th>
<th>Packages</th>
<th>No.</th>
<th>Gross weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4(a)</td>
<td>4(b)</td>
<td></td>
</tr>
</tbody>
</table>

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:

1) The consignment shall not be opened or broken in transit.

2) This pass shall remain in force upto and including ... 19...

— Seal —

Place:

Signature and designation of the Officer issuing the pass.

* Name and full address of the importer.

+ Name and full address of the exporter.

FORM — B

[See rule 16(2)]

Pass for export inter-state of medicine(s) containing opium

(Countfoil)

(Serial No. 19)

Shri/Smt./Kum./M/s. * .......... is/are hereby authorised to export inter-state the undermentioned medicines containing opium to Shri/Smt./Kum./M/s. + ....

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be exported</th>
<th>Total quantity of opium contained in the medicines to be exported</th>
<th>Packages</th>
<th>No.</th>
<th>Gross weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4(a)</td>
<td>4(b)</td>
<td></td>
</tr>
</tbody>
</table>

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:

1) The consignment shall not be opened or broken in transit.

2) This pass shall remain in force upto and including ... 19...

— Seal —

Place:

Signature and designation of the Office issuing the pass.

* Name and full address of the exporter.

+ Name and full address of the importer.

FORM — A

[See rule 15(2)]

Pass for import inter-State of medicine(s) containing opium

(Quadruplicate)

Serial No.:

Dated:

Shri/Smt./Kum./M/s. * .......... is/are hereby authorised to import inter-State the undermentioned medicines containing opium from Shri/Smt./Kum./M/s. + .......... 

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be imported</th>
<th>Total quantity of opium contained in the medicines to be exported</th>
<th>Packages</th>
<th>No.</th>
<th>Gross weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4(a)</td>
<td>4(b)</td>
<td></td>
</tr>
</tbody>
</table>

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:

1) The consignment shall not be opened or broken in transit.

2) This pass shall remain in force upto and including ... 19...

— Seal —

Place:

Signature and designation of the Officer issuing the pass.

* Name and full address of the importer.

+ Name and full address of the exporter.

FORM — A

[See rule 15(2)]

Pass for import inter-State of medicine(s) containing opium

(Duplicate)

(Serial No. 19)

Shri/Smt./Kum./M/s. * .......... is/are hereby authorised to export inter-state the undermentioned medicines containing opium to Shri/Smt./Kum./M/s. + ....

<table>
<thead>
<tr>
<th>Exact description of the medicines</th>
<th>Total quantity of the medicines to be exported</th>
<th>Total quantity of opium contained in the medicines to be exported</th>
<th>Packages</th>
<th>No.</th>
<th>Gross weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4(a)</td>
<td>4(b)</td>
<td></td>
</tr>
</tbody>
</table>

This pass is granted under and subject to the provisions of the Narcotics Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:

1) The consignment shall not be opened or broken in transit.

2) This pass shall remain in force upto and including ... 19...

— Seal —

Place:

Signature and designation of the Officer issuing the pass.

* Name and full address of the exporter.

+ Name and full address of the importer.
FORM — B
[See rule 16(2)]
Pass for export inter-state of medicine(s) containing opium
(Triplicate)
(To be forwarded to the Excise Authority of the
place of import)
Serial No. Dated: 19
Shri/Smt./Kum/M/s. * ... is/are hereby authorised to export
inter-state the undermentioned medicines containing opium
to Shri/Smt./Kum/M/s. + ...

<table>
<thead>
<tr>
<th>Exact description of the medicine</th>
<th>Total quantity of the medicines to be exported</th>
<th>Total quantity of opium contained in the medicines to be exported</th>
<th>Packages</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

This pass is granted under and subject to the provisions of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:

1) The consignment shall not be opened or broken in transit.
2) This pass shall remain in force upto and including ... 19...

Place
Signature and designation of the Officer issuing the pass.

* Name and full address of the exporter.
+ Name and full address of the importer.

FORM — C
[See rule 17(3)]
Pass for transport of opium/medicines containing opium
(Counterfoil)
(For Office use only)
Serial No. Dated: 19
Shri/Smt./Kum/M/s. * ... is/are hereby authorised to transport to + ... from § ... the undermentioned drugs:

<table>
<thead>
<tr>
<th>Exact description of the medicine</th>
<th>Total quantity of the drugs to be transported</th>
<th>Total quantity of opium contained in the medicines containing opium</th>
<th>Packages</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

This pass is granted under and subject to the provisions of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:

1) The consignment shall not be opened or broken in transit.
2) This pass shall remain in force up to an including ... 19...

Place
Signature and designation of the Officer issuing the pass.

* Name of the person or firm authorised.
+ Locality and District of destination of the consignment.
§ Name and full address of the consigner.
© To be filled in when medicines containing opium is to be transported.
**FORM — C**

[See rule 17(3)]

Pass for transport of opium/medicines containing opium (Triplicate)

(To be forwarded to the Inspector of Excise of the place of destination)

<table>
<thead>
<tr>
<th>Serial No.</th>
<th>Dated:</th>
<th>Shri/Smt./Kum/M/s.</th>
<th>is/are hereby authorised to transport to + ... from § ... the undermentioned drugs:—</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>Exact description of the drugs</strong></td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4(a)</td>
</tr>
</tbody>
</table>

This pass is granted under and subject to the provisions of the Narcotics and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder and subject to the following conditions:

1) The consignment shall not be opened or broken in transit.
2) This pass shall remain in force up to an including ... 19 ...

Seal

Place

Signature and designation of the Officer issuing the pass.

* Name of the person or firm authorised.
† Locality and District of destination of the consignment.
§ Name and full address of the consignor.
* To be filled in when medicines containing opium is to be transported.

**FORM — D**

[See rule 24(III)]

Form of Authority

I hereby appoint ... to buy, possess and transport opium on my behalf on my permit No. ...

Date

Identification marks of the agent

1 ...
2 ...

Signature or thumb impression of the person giving the authority.

Approval

Seal

Signature of the Officer authorised to grant the permit.

Place

Date ...

**FORM — E**

[See rule 26(1)]

Name of the Licensee ...

Licence No. ...

Register of Accounts of opium possessed and used during the month of ... 19

<table>
<thead>
<tr>
<th>Opening balance on 1st day of the month of</th>
<th>Quantity received during the month</th>
<th>Total of columns (1) and (3)</th>
<th>Quantity used in preparation of medicines during the month</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4(a)</td>
</tr>
<tr>
<td>Opium</td>
<td>Opium</td>
<td>Opium</td>
<td>Opium</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Closing balance on the last day of the month</th>
<th>Name of medicines in which opium is used and the quantity of such medicines prepared</th>
<th>Quantity received from any other place</th>
<th>Signature of the licensee</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>7</td>
<td>8</td>
<td>9</td>
<td>10</td>
</tr>
<tr>
<td>Opium</td>
<td>Name</td>
<td>Grams</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**FORM — F**

[See rule 26(2)]

Name of Licensee ...

Licence No. ...

Register of Accounts of opium possessed and used during the month of ... 19

<table>
<thead>
<tr>
<th>Date</th>
<th>Opening balance</th>
<th>Quantity received from the depot</th>
<th>Quantity received from any other place</th>
<th>Total of columns (5) and (6)</th>
<th>Quantity used in preparation of medicines</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>Opium</td>
<td>Opium</td>
<td>Opium</td>
<td>Opium</td>
<td>Opium</td>
<td>Opium</td>
</tr>
</tbody>
</table>
The licensee shall not manufacture medicinal opium made on the
name of the Licensee: ... License No ...

Place

FORM - G
(See rule 7)

Name of the Licensee: ... License No ...
Register of daily accounts of medicines containing opium purchased and sold during the month of ... 19

<table>
<thead>
<tr>
<th>Opening balance</th>
<th>Quantity of medicines used and the quantity of such medicines manufactured</th>
<th>Name of medicines</th>
<th>Quantity</th>
<th>Name of medicines</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date</td>
<td>Medicines containing opium</td>
<td>Name of medicines</td>
<td>Quantity</td>
<td>Name of medicines</td>
<td>Quantity</td>
</tr>
<tr>
<td>1</td>
<td></td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>

Quantity of medicines sold
Medicines containing opium
Name of medicines | Quantity | Closing balance Medicines containing opium | Name of medicines | Quantity |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>

FORM - H
(See rule 28)

License for the manufacture of medicinal opium
District of: ---
Number of the Licence: ---
Name and description of the licensee: ---
Residential address: ---
Place of business with boundaries: ---
The person described above, and hereinafter called the licensee is hereby authorised by the Drugs Controller of Goa, Daman and Diu to manufacture medicinal opium from ... to ... subject to the following conditions:

CONDITIONS
(1) The licence is granted on payment of Rupees one thousand only to the licensee personally and is not transferable.
(2) The licensee shall comply with all the provisions of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules framed thereunder.
(3) The licence shall intimate to the Drugs Controller of Goa, Daman and Diu all particulars of the employment or change of his agent (servants, etc.) and pay a fee of Rupees ten only per annum per person. The licensee shall be responsible for the acts and omissions of every person employed by him in carrying on his business and of all his servants, as if the said acts and omissions were his own.
(4) The licensee shall not manufacture medicinal opium save from the materials which he is lawfully entitled to possess.
(5) The licensee shall not manufacture medicinal opium or keep the materials used for the manufacture of medicinal opium at any place except his place of manufacture approved for the purpose.

FORMS
FORM - I

I. Account of the opium used for the manufacture of medicinal opium.

<table>
<thead>
<tr>
<th>Month and date</th>
<th>Quantity in balance</th>
<th>Quantity received</th>
<th>Source of supply</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
</tbody>
</table>

Total of Columns 2 and 3

<table>
<thead>
<tr>
<th>Balance in hand in grams</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>6</td>
</tr>
</tbody>
</table>

FORM - II

II. Account of the medicinal opium manufactured.

<table>
<thead>
<tr>
<th>Month and date</th>
<th>Balance in hand in grams</th>
<th>Quantity manufactured in grams</th>
<th>Total of columns 2 and 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Balance in hand in grams</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>6</td>
</tr>
</tbody>
</table>

Signature with date of the licensee | Signature with date of the Drugs Controller | Remarks if any |
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>10</td>
<td>11</td>
</tr>
</tbody>
</table>
(13) In case of breach of any of the conditions of the licence, the Drugs Controller, may cancel or suspend the licence or in lieu thereof impose a penalty not exceeding ten thousand rupees.

(13) The imposition of a penalty or the cancellation or suspension of this licence under the foregoing condition shall not operate as a bar to prosecution for any offences which may have been committed under the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or the rules, regulations or orders made thereunder from time to time.

(14) If the licensee has in his possession on the expiry, cancellation, or suspension of this licence, any stock of the drug, he shall deliver it up to the Drugs Controller.

(15) The licensee shall not manufacture medicinal opium in a quantity larger than his requirements for one month.

Dated the .. . day of ... 19 ...  

Seal:  

Place:  

Drugs Controller  

Goa, Daman and Diu  

--- FORM - NDPS - I  

[Licence is hereby granted to ... of ... following the profession of ... at ... (hereinafter called the licensee authorising him under and subject to the provisions of the Narcotic Drugs and Psychotropic Act, 1985 (Central Act 61 of 1985) and the rules made thereunder.

(a) to possess and sell, otherwise than on prescription, manufactured drugs (other than prepared opium and coca leaf); and

(b) to manufacture the manufactured drugs or any preparations containing manufacture from the materials which he is lawfully entitled to possess at his shop situated at ... in the district of ... during the period commencing on ... and ending on ... on payment of a fee ... of Rs. ... (Rupees only) and subject to the conditions hereinafter mentioned, namely:—

1. The licensee shall purchase all the manufactured drugs to be sold under this licence from a dealer in manufactured drugs licensed under the Goa, Daman and Diu Narcotic Drugs any Psychotropic Substances Rules, 1987, or under the corresponding rules for the time being in force in any part of India, or in accordance with condition 8 or import in the State such drugs from abroad in accordance with the rules issued under section 8 of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985), by the Central Government. He shall not receive or have in his possession the manufactured drugs, obtained otherwise than as permitted under this condition. Nor shall he receive or have in his possession any quantity of—

(a) coca derivatives containing in the aggregate more than * of cocaine.

(b) Opium derivatives containing in the aggregate more than * of either morphine diacetylmorphine or both.

(c) medicinal hemp exceeding * in the case of extract and * in the case of tinctures.

In case of preparations which are admixtures of coca derivatives and opium derivatives, the limit shall be fixed with reference to the cocaine and morphine contents respectively and not with reference to the quantity or bulk of the preparations, and the bottles, phial, packages or other containers of the preparations or labels affixed to them shall exhibit the statement of the quantity of the manufactured drugs present in each container or sufficient particulars to admit of the ready calculation of such quantity.

2. The licencee shall not keep, store or sell manufactured drugs in any place except in his shop described above.

If he wishes to remove any manufactured drugs from one place to another he shall first obtain a transport authorisation from the Licensing Authority for the purpose.

3. The Licensee shall be responsible for the acts and omissions of every person employed by him for carrying on the business. If the act and omissions of all his servants shall be construed as if the said acts and omissions were his own.

4. The Licensee shall not sell any manufactured drug except

(a) to a dealer in manufactured drugs or a chemist or an approved practitioner licensed under the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987, or under the corresponding rules for the time being in force in any part of India outside the Union territory of Goa, Daman and Diu in accordance with the provisions of rule 32 of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987.

(b) to an approved practitioner to the extent that he is permitted to possess it for use in his practice according to the provisions of rule 32 of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987.

(c) to a person especially authorised by the Licensing Authority to possess and use it under rule 55 of Goan, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987.

(d) to a person holding an export authorisation granted under rule 56 of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987 or under the rules made by the Central Government under section 9 of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985).

(e) to Medical Officer-in-charge of a Government grant-aided medical institution or hospital or dispensary or a Registered Medical practitioner of a municipal dispensary or in charge of hospital or dispensary belonging to a Mission or any other corporate body or to a Medical Officer-in-charge of a hospital or dispensary belonging to a railway or private nursing homes, registered with the Directorate of Health Services, Panaji-Goa.

(f) to a Government Officer who is duly authorised to possess it under rule 53 of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules 1987 provided that a receipt is obtained by the Licensee from such Officer and kept on his record:

Provided that:—

(i) the quantity of the drug sold shall not exceed the quantity which such dealer, chemist, practitioner, person or Officer, as the case may be, may lawfully possess:

(ii) the drug shall not be delivered to any person not licensed or otherwise authorised to be in possession of the drug, who purports to receive it on behalf of a person so licensed or otherwise authorised by the person producing an authority in writing signed by the said so licensed or authorised to receive the drugs on his behalf and unless the licensee is satisfied that the authority is genuine, and

(iii) Where, for the removal of the drug on authorisation is required, the Licensee shall, before permitting the removal of the drug from his premises by the purchaser, satisfy himself that the purchaser has obtained the necessary authorization from the Licensing Authority.

5. The Licensee may import, in the Union territory, export from the Union territory or transport the manufactured drugs by rail or sea subject to the following conditions:—

(a) The parcel of the manufactured drug when sent by post, shall be sent by registered parcel only;

(b) The parcel, whether sent by rail or by post, shall be insured;

(c) The parcel shall be covered by an authorization issued by the competent authority at the place to which the parcel is addressed;

(d) The parcel shall be accompanied by a declaration showing the names of the consignor and the consignee,
the contents of the parcel in detail, the number and description of the various drugs, the quantity of each, and the date of purchase or sale, and the name and address of the person or firm from whom the purchase was made or to whom or on whose behalf the drug was sold, as the case may be.

(2) The licensee shall preserve the said accounts and authorise the same to be inspected by any such officer as the Licensing Authority may from time to time designate, at any time after entry in the account book has been made, or in respect of any operation, entry or omission provided for under the said rules.

(3) The licensee shall furnish to the Drug Controller’s Office or any other officer duly authorised by him in his behalf, within a month after the end of each quarter of the year, the information regarding the purchase, sale and consumption of manufactured drugs during the preceding quarter and the stocks of manufactured drugs held by him in balance on the last day of the quarter in form prescribed by the Licensing Authority for the purpose.

7. (1) This licence may be cancelled or suspended by the Licensing Authority at any time—

(a) for non-payment of fee payable by the licensee;

(b) for default or violation by himself or by any servant or person acting on his behalf of any of the conditions specified in his licence or of the provisions of the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987.

(c) if the Licensee is convicted of a breach of peace or of any offence under the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under any law for the time being in force relating to arms, revenue or of any other criminal offences during the currency of the licence;

(d) if the Licensee violates any of the condition imposed on him by the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under any law for the time being in force relating to arms, revenue or of any other criminal offences;

(e) if the Licensee violates any of the condition imposed on him by the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under any law for the time being in force relating to arms, revenue or of any other criminal offences;

(f) if the Licensee violates any of the condition imposed on him by the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under any law for the time being in force relating to arms, revenue or of any other criminal offences;

(g) if the Licensee violates any of the condition imposed on him by the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under any law for the time being in force relating to arms, revenue or of any other criminal offences;

(h) if the Licensee violates any of the condition imposed on him by the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under any law for the time being in force relating to arms, revenue or of any other criminal offences;

(i) after giving the Licensee 15 days notice, or if the Licensee desires to surrender his licence within 15 days from the receipt of such notice from him.

(2) When such licence is cancelled, suspended or surrendered the Licensee shall forthwith make over to the Licensing Authority or to such other Officer as he may appoint, his licence together with all manufactured drugs in his possession.

(3) The Licensee shall be bound to purchase in such quantity not exceeding that which he is likely to sell in one month and at such rates as the Licensing Authority may direct, any manufactured drugs that may be delivered to the Licensing Authority by any other licensee whose licence has expired or has been cancelled or suspended.

(9) All preparations containing not more than 0.1 per cent of cocaine or 0.2 per cent of morphine and any preparations declared by the Central Government by notification in the Gazette of India in pursuance of finding under article 8 of the Geneva Convention, not to be a manufactured drugs, may be imported, transported, possessed, and sold without restriction.

FORM—N.D.P.S.3

[See rule 29(21)]

Licence for the possession and sale, on prescription of manufactured drugs by Chemist and Medical Practitioners.

No. ____________ Date: ____________

Licence is hereby granted to ... following the profession of ... at ... (hereinafter called the Licencee) authorising him under and subject to the provisions of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and the rules made thereunder to possess, sell or dispense, on prescription only, manufactured drugs at his shopdispensary situated at ... in the district of ... during the period commencing on ... and ending on ... on payment of a fee of Rs. (Rupees ... only) subject to the conditions hereinafter mentioned, namely:

1. The licensee shall purchase all manufactured drugs to be sold or dispensed under this licence, from a dealer in manufactured drug licensed under the Goa, Daman and Diu Narcotic Drugs and Psychotropic Substances Rules, 1987 or under the corresponding rules for the time being in force in any part of India, or in accordance with condition.

He shall not receive or have in his possession any manufactured drugs which are not specified in this condition or which have been obtained otherwise than as permitted under this condition, nor shall he possess them in quantities exceeding those specified below:

(a) cocoa derivatives containing in the aggregate more than ... of cocaine.

(b) opium derivatives containing in the aggregate more than ... of either morphine, diacetylmorphine or both.

(c) medicinal hemp exceeding ... in case of extract and ... in case of tincture.

In the case of preparations and admixtures of cocoa derivatives and opium derivatives, the limit shall be fixed with reference to the cocaine and morphine contents, and not with reference to the quantity or bulk of the preparation and the bottle phials or other containers of these preparations or labels affixed to them shall plainly exhibit the actual quantity of the manufactured drugs present in each container or sufficient particulars to admit of the ready calculation of such quantity.

2. (a) The Licensee, unless he is a Registered Medical Practitioner shall not keep, store, sell or dispense the manufactured drugs in any place except in his dispensary described above.

(b) If the Licensee, is a Registered Medical Practitioner, he may carry with him, from place to place manufactured drugs in quantities not exceeding those specified in conditions 1 above.

3. The Licensee shall be responsible for the acts of commissions and omissions of every person, employed by him in carrying on the business of the said dispensary, and of all his servants and agents as if the said acts of commission and omissions were his own.

4. (1) The Licensee shall not sell or dispense any manufactured drugs except on a bonafide prescription given by himself, if he is a Registered Medical Practitioner, or by any other Registered Medical Practitioner not in larger quantity nor to any other person other than may be specified in the prescription, provided the prescription is not given for the use of the prescribed himself.

(2) A prescription for the supply of any manufactured drugs must comply with the following conditions:

(a) The prescription shall be in writing, and shall be signed and signed by a Registered Medical Practitioner with his full name, qualifications, and address and shall also specify the name and address of the person to whom it is given and the total quantity of the drugs to be supplied thereof.

(b) The prescription shall not be given for the use of the prescriber himself.
(c) A prescription given by a Registered Dental Practitioner shall be only for the purpose of dental treatment of and shall be marked 'For Local Dental Treatment Only'.

(3) When cocoa derivatives are to be sold or dispensed, the Licensee shall ensure that the prescription is marked with the words 'Not to be Repeated' and shall not supply cocoa derivatives on the same prescription except in pursuance of fresh directions on the prescription by the Registered Medical Practitioner by whom it was originally issued dated and signed with his name in full.

(4) Where opium derivatives or medicinal hemp are to be sold or dispensed:

(a) if the prescription does not bear a superscription by a Registered Medical Practitioner stating that it is to be repeated and what interval of time it is to be repeated and how many times it is to be repeated, the Licensee shall sell the drugs once only on such prescription, and shall retain the prescription, provided that he shall first warn the person presenting the prescription that, unless it bears such a superscription as aforesaid it will be retained.

(b) if the prescription bears a superscription as aforesaid, and if it appears that opium derivatives or medicinal hemp are already sold on the prescription six times, or such number of times as the prescription is required to be repeated, or that the interval specified in the superscription has been elapsed since the prescription was last dispensed, he shall not sell the drugs on such prescription, unless it is further superscribed in that behalf by the Registered Medical Practitioner.

5. The Licensee shall mark on every prescription dispensed by him, his name, the address of the premises at which and the date on which it was dispensed. In the case of a prescription made upon a prescription which contains manufactured drugs, the bottle or other receptacle or the wrapper or other covering in which such preparation is enclosed shall bear clearly marked upon it the amount and percentage of cocaine or morphine or diacetylmorphine or medicinal hemp contained in such preparation, provided that if the preparation in the form of uniformly divided dosal units e.g., pills, powders, tablets, capsules, etc, it shall be sufficient if the bottle or other receptacle or the wrapper or other covering in which such preparation is enclosed bears clearly marked upon it the amount and percentage of cocaine or morphine contained in each such dosal unit.

6. Where the prescription has to be returned to the person who presents it, the Licensee shall, on the first sale thereof, take and keep a copy, of it and on the occasions of each subsequent sale, not thereon, the date of the sale and also sign and seal it.

5. The Licensee may import, export or transport manufactured drugs by rail or inland post subject to the following conditions:

(a) the parcel of manufactured drugs when sent by post, shall be sent by registered parcel only.

(b) the parcel shall be covered by an authorization issued by competent authority at the place to which the parcel is addressed;

(c) the parcel shall be accompanied by a declaration showing the names of the consignor and the consignee, the contents of the parcel in detail, the number and date of authorization covering the import, export or transit as the case may be, and the number of the licence, if any, held by the consignor and the consignee.

6. The Licensee shall file and preserve for two years all prescriptions upon which manufactured drugs have been sold or dispensed by him, and shall produce such prescription alongwith this licence and any manufactured drugs that may be in his possession for inspection on demand by the Licensing Authority duly authorised by him.

The Licensee shall maintain a register in such form as may be approved by the Licensing Authority, wherein he shall from time to time, record in respect of the manufactured drugs dispensed by him the full names and addresses of the Registered Medical Practitioners prescribing drugs. The Licensee shall similarly record in the said register a true account of the kind and quantity of the manufactured drugs dispensed and the balance held by him in stock. The Licensee shall, before the seventh day of each calendar month, furnish to the Drugs Controller's Office as he may appoint in this behalf a copy of the entries made by him in the register during the preceding calendar month.

7. (i) This Licence may be cancelled or suspended by the Licensing Authority at any time.

(a) for non-payment of duty or fee payable by the Licensee.

(b) for default or violation by himself or by any servant or person acting on his behalf, of any of the conditions specified in the licence or if the provisions of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985).

(c) if the Licensee is convicted of any offence under the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under the law for the time being in force relating to excise revenue or a breach of the peace or of any other criminal offence during the currency of the licence.

(d) if the Licensee violates any of the conditions imposed on him by the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) or under the rules, regulations or order made thereunder.

(e) after giving the Licensee fifteen days notice, or if the Licensee desires to surrender his licence, within fifteen days from the receipt of such notice from him.

(ii) When such licence is cancelled, suspended or surrendered, the Licensee shall forthwith make over to the Licensing Authority or such other officer as he may appoint, the licence together with all the manufactured drugs in his possession.

8. The Licensee shall be bound to purchase in such quantity not exceeding that which he is likely to sell in two months and at such rates as the Licensing Authority may direct, any manufactured drugs, that may be delivered to the Licensing Authority by any other Licensees whose licence has expired or has been cancelled or suspended.

9. All preparations containing not more than 0.1 per cent of cocaine or 0.2 percent of morphine and any preparations declared by the Central Government by notification in the Gazette of India in pursuance of finding under article 8 of the Geneva Convention, not to be a manufactured drugs, may be imported, transported, possessed and sold without restriction.

Granted this the ... day of ...

— Seal —

Address : Licensing Authority

* To be fixed by the Licensing Authority.

FORM — NDPS — 3

[See rule 39(2)]

Authorization No. Dated : Special Authorization for Registered Medical Practitioners for the possession of coca derivatives for the use in the exercise of their profession but not for sale.

Dr. ..... is hereby authorised to possess coca derivatives containing not more than ... grams of cocaine for the use in the exercise of his profession in his dispensary situated at ... Taluka of ... of the District of .... This Authorization which is granted free of charge, will be valid until the 31st March, 19... and is issued subject to the following conditions:

1. That the Authorization Holder shall be bound by the provisions of the Narcotic Drugs and Psychotropic Substances Act, 1985 (Central Act 61 of 1985) and any
FORM — NDPS 4

(See rule 58)

Authorization for the inter-provincial Import of Manufactured drugs into the Union Territory of Goa, Daman and Diu

PART I

(To remain in the Office of issue)

Shri/Smt/Kum/M/s. * ... hereby authorised to import are

the undermentioned drugs from Shri/Smt/Kum/M/s. **...

<table>
<thead>
<tr>
<th>Exact description of the drugs</th>
<th>Total quantity of the drugs to be imported</th>
<th>Percentage of the drugs contents</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This Authorization must be used within two months from the date of its issue.

The bulk of the consignment shall not be opened or broken in transit.

Dated the ... 19

Place

Seal

Licensing Authority

* Name and full address of the importer.
** Name and full address of the exporter.
Authorization for the Inter-Provincial Export of Manufactured Drugs from the Union territory of Goa, Daman and Diu

PART I

(To be retained in the Office of issue)

is Shri/Smt/Kum/M/s. * ... hereby authorized to export are the undermentioned drugs to Shri/Smt/Kum/M/s ** ...

by + ... in $ ...  

<table>
<thead>
<tr>
<th>Exact description of the drugs</th>
<th>Total quantity of the drugs to be exported</th>
<th>Percentage of the drugs contents</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This authorization will remain in force upto the ... day of the month of ... 19

The bulk of the consignment shall not be opened or broken in transit.

Dated ... 19

Place:

Seal

Licensing Authority

* Name and full address of consigner.

** Name and full address of consignee.

+ Route and mode of conveyance.

§ Number and description of packages.

---

Authorization for the Inter-Provincial Export of Manufactured Drugs from the Union territory of Goa, Daman and Diu

PART II

(To be handed over to the consignee to accompany the consignment)

is Shri/Smt/Kum/M/s. * ... hereby authorized to export are the undermentioned drugs to Shri/Smt/Kum/M/s ** ...

by + ... in $ ...  

<table>
<thead>
<tr>
<th>Exact description of the drugs</th>
<th>Total quantity of the drugs to be exported</th>
<th>Percentage of the drugs contents</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This authorization will remain in force upto the ... day of the month of ... 19

The bulk of the consignment shall not be opened or broken in transit.

Dated ... 19

Place:

Seal

Licensing Authority

* Name and full address of consigner.

** Name and full address of consignee.

+ Route and mode of conveyance.

§ Number and description of packages.

---

Authorization for the Inter-Provincial Export of Manufactured Drugs from the Union territory of Goa, Daman and Diu

PART IV

(To be issued in the case of despatch of drugs by parcel by sea and handed over to the consigner for production at the Port Office of despatch)

is Shri/Smt/Kum/M/s. * ... hereby authorized to export are the undermentioned drugs to Shri/Smt/Kum/M/s ** ...

by + ... in $ ...  

<table>
<thead>
<tr>
<th>Exact description of the drugs</th>
<th>Total quantity of the drugs to be exported</th>
<th>Percentage of the drugs contents</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**FORM — NDPS 6**

(See rule 60)

**Authorization for the transport of manufactured drugs within the Union territory of Goa, Daman and Diu**

**DUPLICATE**

(To be forwarded to the authority of the place from which the drugs are to be transported)

<table>
<thead>
<tr>
<th>Exact description of the drugs</th>
<th>Total quantity of the drugs to be exported</th>
<th>Percentage of the drugs contents</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This authorisation will remain in force upto the ... day of the month of ... 19

The bulk of the consignment shall not be opened or broken in transit.

Dated ... 19

Place:

Licensing Authority

* Name and full address of consigner.

** Name and full address of consignee.

+ Route and mode of conveyance.

$ Number and description of packages.

---

**FORM — NDPS 6**

(See rule 60)

Authorization for the transport of manufactured drugs within the Union territory of Goa, Daman and Diu

**COUNTERFOIL**

(To remain attached to book)

is Shri/Smt/Kum/M/s. * ... hereby authorised to transport to ** ... from X ... the undermentioned drugs:—

<table>
<thead>
<tr>
<th>Exact description of the drugs</th>
<th>Total quantity of the drugs to be transported</th>
<th>Percentage of the drugs contents</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This authorisation must be used within two months from the date of its issue.

Dated ... 19

Place:

Licensing Authority

* Name of the person or firm authorised.

** Locality and District of destination of the consignment.

X Name and full address of the firm supplying the drugs.

+ Route and mode of conveyance.

$ Number and description of packages.

---

**FORM — NDPS 6**

(See rule 60)

Authorization for the transport of manufactured drugs within the Union territory of Goa, Daman and Diu

**TRIPLICATE**

(To be handed over to the applicant to accompany the consignment)

<table>
<thead>
<tr>
<th>Exact description of the drugs</th>
<th>Total quantity of the drugs to be transported</th>
<th>Percentage of the drugs contents</th>
<th>Remarks if any</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

This Authorization must be used within two months from the date of the issue.

Dated ... 19

Place:

Licensing Authority

* Name of the person or firm authorised.

** Locality and District of destination of the consignment.

X Name and full address of the firm supplying the drugs.

+ Route and mode of conveyance.

$ Number and description of packages.

---

* Name of the person or firm authorised.

** Locality and District of destination of the consignment.

X Name and full address of the firm supplying the drugs.
This Authorization must be used within two months from the date of its issue.

Dated ... 19
Place:
Seal

Licensing Authority

This authorization is to remain in force up to ... day of the month of ... 19
The drug covered by it shall be conveyed intact by ...
In $ ...
Dated ... 19
Place:
Seal

Licensing Authority

* Name of the person or firm authorised.
** Locality and District of destination of the consignment.
X Name and full address of the firm supplying the drugs.
+ Route and mode of conveyance.
$ Number and description of packages.

** Form of Accounts of coca derivatives to be kept by Registered Medical Practitioners who have been granted Special authorization for the possession of coca derivatives for use in the exercise of their profession.**

<table>
<thead>
<tr>
<th>Month</th>
<th>19</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cocaine derivatives powder</td>
<td></td>
</tr>
<tr>
<td>Cocaine derivatives tabloids</td>
<td></td>
</tr>
<tr>
<td>Cocaine derivatives solution</td>
<td></td>
</tr>
<tr>
<td>Remarks</td>
<td></td>
</tr>
<tr>
<td>Grams</td>
<td>Milligrams</td>
</tr>
</tbody>
</table>

Opening Balance

Purchased during the month ...

Total ...

Used during the months

Closing Balance

Carried to next month.

By order and in the name of the Lieutenant Governor of Goa, Daman and Diu.

L. J. Menezes Pais, Under Secretary (Health).