

File No: BIO/CT/19/000083  
Government of India  
Directorate General of Health Services  
Central Drugs Standard Control Organization  
(Biological Division)

From:

The Drugs Controller General, India  
Directorate General of Health Services,

FDA Bhawan Kotla Road,  
New Delhi-110002

To,

M/s Intas Pharmaceuticals Limited,  
Plot no. 423/P/A, Sarkhej-Bavla Highway, Village: Moraiya,  
Taluka: Sanand, Ahmedabad-382213, Gujarat, India

Subject: Application for grant of permission to conduct clinical trial entitled – “A Randomized, Assessor-Blind, Multiple-Dose, Two-Treatment, Two-Period, Two-Way Crossover Study Comparing The Pharmacodynamics of INTP1 From Intas Pharmaceuticals Ltd., India And Neupogen of Amgen Inc., USA following Subcutaneous Injections In Healthy, Adult Human Subjects vide Protocol No: 0365-19; Version: 1.0; Date: 07 August 2019” for export purpose only - regarding

Ref.: Your Application No. BIO/CT04/FF/2019/16716 dated 02-Oct-2019

Sir,

With reference to your Application No.: BIO/CT04/FF/2019/16716 dated 02-Oct-2019, please find enclosed herewith the permission in FORM CT-06 for conduct of subject clinical trial for export purpose only under the provisions of New Drugs and Clinical Trial Rules, 2019.

The permission granted by the Central Licensing Authority to conduct clinical trial under this Chapter shall be subject to following conditions, namely:

- (I) Clinical trial at each site shall be initiated after approval of the clinical trial protocol and other related documents by the Ethics Committee of that site, registered with the Central Licensing Authority under rule 8;
- (II) Where a clinical trial site does not have its own Ethics Committee, clinical trial at that site may be initiated after obtaining approval of the protocol from the Ethics Committee of another trial site; or an independent Ethics Committee for clinical trial constituted in accordance with the provisions of rule 7:  
Provided that the approving Ethics Committee for clinical trial shall in such case be responsible for the study at the trial site or the centre, as the case may be:  
Provided further that the approving Ethics Committee and the clinical trial site or the bioavailability and bioequivalence centre, as the case may be, shall be located within the same city or within a radius of 50 kms of the clinical trial site;
- (III) In case an ethics committee of a clinical trial site rejects the approval of the protocol, the details of the same shall be submitted to the Central Licensing Authority prior to seeking approval of another Ethics Committee for the protocol for conduct of the clinical trial at the same site;
- (IV) The Central Licensing Authority shall be informed about the approval granted by the Ethics Committee within a period of fifteen working days of the grant of such approval;
- (V) Clinical trial shall be registered with the Clinical Trial Registry of India maintained by the Indian Council of Medical Research before enrolling the first subject for the trial;



FORM CT-06

(See rules 22, 25, 26, 29 and 30)

PERMISSION TO CONDUCT CLINICAL TRIAL OF NEW DRUG OR  
INVESTIGATIONAL NEW DRUG

The Central Licencing Authority hereby permits M/s Intas Pharmaceuticals Limited (Biopharma Division), Plot no. 423/P/A, Sarkhej-Bavla Highway, Village: Moraiya, Taluka: Sanand, Ahmedabad-382213, Gujarat, India to conduct clinical trial of the new drug or investigational new drug as per Protocol No: 0365-19; Version: 1.0; Date: 07 August 2019 for export purpose only in the below mentioned clinical trial sites.

2. Details of new drug and clinical trial site [as per Annexure].

3. This permission is subject to the conditions prescribed in part A of Chapter V of the New Drugs and Clinical Trials Rules, 2019 under the Drugs and Cosmetics Act, 1940.

Place: New Delhi

Date:

V G SOMANI  
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DN: cn=CENTRAL DRUGS STANDARD  
CONTROL ORGANIZATION, o=V G SOMANI  
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(Dr. V. G. Somani)  
Central Licencing Authority

**Annexure:****Details of new drug or investigational new drug:**

Names of the new drug or investigational new drug:	Granulocyte Colony Stimulating Factor (Filgrastim)
Therapeutic class:	Cytokines
Dosage form:	Solution for Injection, Pre-filled syringe
Composition:	Each 0.8 mL contains 480 mcg of Filgrastim
Indications:	Filgrastim is indicated for: (1) Cancer patients receiving myelosuppressive chemotherapy, (2) Patients with acute myeloid leukemia receiving induction or consolidation chemotherapy, (3) Cancer patients receiving bone marrow transplant, (4) Patients undergoing peripheral blood progenitor cell collection and therapy, (5) Patients with severe chronic neutropenia, (6) Patients acutely exposed to myelosuppressive doses of radiation (hematopoietic syndrome of acute radiation syndrome), (7) Patients with HIV infection

**Details of clinical trial site:**

S. No.	Name and Address of Clinical Trial Site	Ethics Committee Details	Name of Principal Investigator
1	Lambda Therapeutic Research Ltd., Lambda house, Plot No.38, Survey no. 388, Near Silver Oak Club, S.G. Highway, Gota, Ahmedabad-382481, Gujarat, India.	Riddhi Medical Nursing Home, Institutional Ethics Committee (RMNH IEC) A/101, Jalaram Plaza, Jawahar Chowk, Maninagar, Ahmedabad-380008, Gujarat, India EC Reg. No.: ECR/886/Inst/GJ/2016	Dr. Shri Krishna Kolte