A NAME OF THE MEDICINAL PRODUCT

LONOUEX PRE-FILLED SYRINGE 6MG/0.6ML

## QUALITATIVE AND QUANTITATIVE COMPOSITION

Each pre-filled syringe contains 6 mg of lineafilgrastim\* in 0.6 ml solution Each ml of solution for injection contains 10 mg of lipeefilerastim.

The active substance is a covalent conjugate of filgrastim\*\* with methoxy polyethylene glycol (PEG) via a carbohydrate linker.

This is based on protein content only. The concentration is 20.9 mg/ml (i.e. 12.6 mg per pre-filled syringe) if the PEG moiety and the carbohydrate linker are included. \*\*Filerastim (recombinant methionyl human granulocyte-colony stimulating factor (G\_CSEI) is produced in Escherichia coli cells by recombinant DNA technology

The potency of this medicinal product should not be compared to the potency of another negulated or non-negulated protein of the same therapeutic class. For more information, see section 5.1.

# Excipients with known effect Each pre-filled syringe contains 30 mg sorbitol.

For the full list of excinients, see section 6.1



Solution for injection in pre-filled syringe (injection) Clear colourless solution

A CLINICAL PARTICULARS

### 4.1 Therapeutic indications

Longuex is indicated in adults for reduction in the duration of neutropenia and the incidence of febrile neutropenia in patients treated with cytotoxic chemotherapy for malignancy (with the exception of chronic myeloid leukaemia and myelodysolastic syndromes).

## 4.2 Posology and method of administration

Lonquex treatment should be initiated and supervised by physicians experienced in oncology or haematology.

One 6 mg dose of lipegfilgrastim (a single pre-filled syringe of Lonquex) is recomber a characteristic control of the control chemotherapy.

## Special populations

In clinical studies with a limited number of elderly nationss there was no relevant an clinical studies with a limited fidinois of electry patents, there was no relevant age-related difference with regard to the efficacy or safety profiles of lipegfilgrastim Therefore, no adjustment of the dose is necessary for elderly patients.

Patients with renal impairment
Currently available data are described in section 5.2, but no recommendation on a nosology can be made

### Patients with benatic impairment

Currently available data are described in section 5.2, but no recommendation on a nosology can be made

Paediatric population
The safety and efficacy of Lonquex in children and adolescents aged up to 17 years have not yet been established. Currently available data are described in sections 4.8. 5.1 and 5.2

## Method of administration

The solution is injected subcutaneously (SC). The injections should be given into the abdomen, upper arm or thigh.

Self-administration of Lonquex should only be performed by patients who are well motivated, adequately trained and have access to expert advice. The first injection should be performed

under direct medical supervision For instructions on handling of the medicinal product before administration, see section 6.6.

## 4.3 Contraindications

## Hypersensitivity to the active substance or to any of the excipients listed in section 6.1

### 4.4 Special warnings and precautions for use Traceability

ornve the traceability of biological medicinal products, the trade name and batch number of the administered medicinal product should be clearly recorded in the

General The safety and efficacy of Longuex have not been investigated in patients receiving

# high dose chemotherapy. Lonquex should not be used to increase the dose of cytotox chemotherapy beyond established dose regimens.

Allergic reactions and immunogenicity
Patients who are hypersensitive to G-CSF or derivatives are also at risk of hypersensitivity reactions to lipegfligrastim due to possible cross-reactivity. No lipegfligrastim therapy should be commenced in these patients because of the risk of

Most biological medicinal products elicit some level of anti-drug antibody response. This

antibody response can, in some cases, lead to undesirable effects or loss of efficacy. If a patient fails to respond to treatment, the patient should undergo further evaluation. If a serious allergic reaction occurs, appropriate therapy with close patient follow-up over several days should be administered.

Haematopoietic system
Treatment with lipegfilgrastim does not preclude thrombocytopenia and anaemia caused by myelosuppressive chemotherapy. Lipegfilgrastim may also cause reversible thrombocytopenia (see section 4.8). Regular monitoring of the platelet count and haematorit is recommended. Special care should be taken when administering single or combination chemotheraneutic medicinal products that are known to cause seve thrombocytopenia.

Leukocytosis may occur (see section 4.8). No adverse events directly attributable to leukocytosis have been reported. Elevation in white blood cells (WBĆ) is consistent with the pharmacodynamic effects of lipegfilgrastim. A WBC count should be performed at regular intervals during therapy owing to the clinical effects of lipegfilgrastim and the potential for leukocytosis. If WBC counts exceed 50 x 10°/l after the expected nadir, lineefilerastim should be discontinued immediately.

Increased haematopoietic activity of the bone marrow in response to growth factor therapy has been associated with transient positive bone-imaging findings. This should be considered when interpreting bone-imaging results.

### Patients with myeloid leukaemia or myelodysplastic syndromes

Granulocyte-colony stimulating factor can promote growth of myeloid cells and some non-myeloid cells in vitro.

The safety and efficacy of Longuex have not been investigated in patients with chronic myeloid leukaemia, myelodysplastic syndromes or secondary acute myeloid leukaemia; it should therefore not be used in such patients. Particular care should be taken to distinguish the diagnosis of blast transformation of chronic musloid loukagenia from

### Solenic adverse reactions

poleriik, auverse reactions penerally asymptomatic cases of splenomegaly have been reported after administration of lipegfligrastim (see section 4.8) and infrequent cases of splenic rupture, including fatal cases, have been reported after administration of G-CSF or derivatives (see section 4.8). Spleen size should therefore be carefully monitored (e.g. clinical examination, ultrasound). A diagnosis of splenic rupture should be considered in patients reporting left upper abdominal pain or shoulder tip pain.

### Pulmonary adverse reactions

Pulmonary adverse reactions, Pulmonary adverse reactions, in particular interstitial pneumonia, have been reported after administration of lipegfilgrastim (see section 4.8). Patients with a recent history of pulmonary infiltrates or pneumonia may be at higher risk.

The onset of nulmonary symptoms such as cough, fever and dyspance in association with The crise or pulmonary symptoms such as cough, rever and dyspinose in a sociation with radiological signs of pulmonary infiltrates and deterioration in pulmonary function together with an increased neutrophil count may be preliminary signs of Acute Respiratory Distress Syndrome (ARDS) (see section 4.8). In such circumstances Lonquex should be discontinued at the discretion of the physician and appropriate treatment given.

## Vascular adverse reactions

Capillary leak syndrome has been reported after administration of G.CSE or derivatives Lapillary leak syndrome has been reported after administration of U-SF or derivatives and is characterised by hypotension, hypoalbuminaemia, oedema and haemoconcentration. Patients who develop symptoms of capillary leak syndrome should be closely monitored and receive standard symptomatic treatment, which may include a need for intensive care (see section 4.6).

Patients with sickle cell anaemia
Sickle cell crisis has been associated with the use of G-CSE or derivatives in natients Sickie cell drisis has been associated with the use of G-CSF or derivatives in patients with sickie cell anaemia (see section 4.9). Physicians should therefore severice caution when administering Lonquex in patients with sickle cell anaemia, monitor appropriate clinical parameters and laboratory results and be attentive to the possible association of lipegifigastim with splenic enlargement and vaso-occlusive crisis.

Activitis has been eported after C-CSF administration in healthy subjects and in cancer

nationts. The symptoms experienced included fever abdominal pain malaise, back pair patients. The symptoms experienced included rever, abudinina pain, inclaise, back pain and increased inflammatory markers (e.g. C-reactive protein and white blood cell count). In most cases aortitis was diagnosed by CT scan and generally resolved after withdrawal of C-CSF. See also section 4.8.

### Hypokalaemia

a may occur (see section 4.8). For nationts with increased risk on pokalaemia due to underling disease or co-medications, it is recommende e serum potassium level carefully and to substitute potassium if necessai Glomerulonenhritis

Glomerulonenhritis has been reported in patients receiving filerastim lennerastim or omerunnepmins has been reported in patients receiving nigrastini, jeriograsi gefligrastim. Generally, events of glomerulonephritis resolved after dose redu ithdrawal of filigrastim, lenograstim or pegfilgrastim. Urinalysis monitoring is commended (see section 4.8).

Excinients with known effect

This medicinal product contains sorbitol. Patients with rare hereditary problems of fructose intolerance should not use this medicinal product.

This medicinal product contains less than 1 mmol sodium (23 mg) per pre-filled syringe, i.e. essentially 'sodium-free'.

## 4.5 Interaction with other medicinal products and other forms of interaction

4.5 miteraction with order meteorization products and other forms of miteraction. Due to the potential sensitivity of ranging dyviding imposition cells to cytotomic order and an administration of cytotoxic chemotherapy. Concomitant use of lipogrigations administration of cytotoxic chemotherapy. Concomitant use of lipogrigations with chemotherapy time decidinal product has not been evaluated in patients. In animal models, concomitant administration of C-CSF and 5-fluorourial (S-FU) or other aritmetabolites has been shown to potentiate myelosuppression.

The safety and efficacy of Lonquex have not been evaluated in patients receiving hemotherapy associated with delayed myelosuppression, e.g. nitrosoureas The notential for interaction with lithium, which also promotes the release of rocking to interaction with minuting which also promotes the release of strophils, has not been specifically investigated. There is no evidence that such an eraction would be harmful.

## 4.6 Fertility, pregnancy and lactation

## Pregnancy

Pregnancy
There are very limited data (less than 300 pregnancy outcomes) on the use of lipegfilgrastim in pregnant women. Animal studies have shown reproductive toxicity (see section 5.3). As a precautionary measure, it is preferable to avoid the use of Longuex during pregnancy.

Breast-feeding
It is unknown whether lipegfligrastim/metabolites are excreted in human milk. A risk to
the breast-fed child cannot be excluded. Breast-feeding should be discontinued during treatment with Longuex.

<u>Fertility</u>

No data are available. Animal studies with G-CSF and derivatives do not indicate harmful effects with respect to fertility (see section 5.3).

## 4.7 Effects on ability to drive and use machines

### I onnuex has no or neeligible influence on the ability to drive and use marbines 4.8 Undesirable effects

# Summary of the safety profile The most frequent undesirable effects are musculoskeletal pain and nausea

Capillary leak syndrome, which can be life-threatening if treatment is delayed, has been reported mostly in cancer patients undergoing chemotherapy after admini G-CSF or derivatives (see section 4.4 and section 4.8). Tabulated list of adverse reactions

The safety of lipegfligrastim has been evaluated based on results from clinical studies including 506 patients and 76 healthy volunteers treated at least once with lipegfligrastim. The adverse reactions listed below in table 1 are classified according to system organ class. Frequency groupings are defined according to the following convention: very common (≥1/10), common (≥1/100 to <1/10), uncommon (≥1/1,000 to <1/100), rare (>1/10,000 to <1/1,000), very rare (<1/10,000), not known (cannot be estimated from

Within each frequency grouning, undesirable effects are presented in order of

Table 1: Adverse reactions			
System organ class	Frequency	Adverse reaction	
Blood and lymphatic system disorders	Common	Thrombocytopenia*	
	Uncommon	Leukocytosis*, Splenomegaly*	
Immune system disorders	Uncommon	Hypersensitivity reactions*	
Metabolism and nutrition disorders	Common	Hypokalaemia*	
Nervous system disorders	Common	Headache	
Vascular disorders	Not known	Capillary leak syndrome* Aortitis*	
Respiratory, thoracic and mediastinal disorders	Common	Haemoptysis	
	Uncommon	Pulmonary adverse reactions*, Pulmonary Haemorrhage	
Gastrointestinal disorders	Very common	Nausea*	
Skin and subcutaneous tissue disorders	Common	Skin reactions*	
	Uncommon	Injection site reactions*	
Musculoskeletal and connective tissue disorders	Very common	Musculoskeletal pain*	
General disorders and administration site conditions	Common	Chest pain	
Investigations	Uncommon	Blood alkaline phosphatase increased*, Blood lactate dehydrogenase increased*	

## \*See section "Description of selected adverse reactions" below

Description of selected adverse reactions
Thrombocytopenia and leukocytosis have been reported (see section 4.4). Splenomegaly, generally asymptomatic, has been reported (see section 4.4). Hypersensitivity reactions such as allergic skin reactions, urticaria, angioedema and

### serious allereic reactions may occur. Hypokalaemia has been reported (see section 4.4).

Pulmonary adverse reactions in particular interstitial pneumonia, have been reported remininary adverse reactions, in particular integration priceduring and continuing not over the sceen section 4.4). These pulmonary adverse reactions may also include pulmon oedema, pulmonary infiltrates, pulmonary fibrosis, respiratory failure or ARDS (see section 4.4).

Nausea was very commonly observed in natients receiving chemotherany Skin reactions such as erythema and rash may occur.

Injection site reactions such as injection site induration and injection site pain may occur The most frequent adverse reactions include musculosteletal pains such as bone pain and myalgia. Musculosteletal pains such as bone pain and myalgia. Musculosteletal paints generally of mild to moderate severity, transient and can be controlled in most patients with standard analgerist. However cases of severe musculosteletal pain (mainly bone pain and back pain) have been reported, including cases that led to hospitalisation.

Reversible, mild to moderate elevations in alkaline phosphatase and lactate dehydrogenase may occur, with no associated clinical effects. Elevations in alkaline phosphatase and lactate dehydrogenase most likely originate from the increase in neutronhils Certain adverse reactions have not yet been observed with lineefilerastim, but are

generally accepted as being attributable to G-CSE and derivative Blood and lymphatic system disorders

Splenic rupture including some fatal cases (see section 4.4)

Sickle cell crisis in patients with sickle cell anaemia (see section 4.4)

Cases of capillary leak syndrome have been reported in postmarketing experience after administration of G-CSF or derivatives. These have generally occurred in patients suffering from advanced malignant diseases, having sepsis, taking multiple

chemotherany medicinal products or undergoing apheresis (see section 4.4). Apritis (see section 4.4)

Skin and subcutaneous tissue disorders

- Acute febrile neutrophilic dermatosis (Sweet's syndrome) Cutaneous vasculitis

Renal and urinary disorders

- Glomerulonephritis (see section 4.4)

Beadistric population
The experience in children is limited to a single-dose phase 1 study in 21 paedistric
patients aged 2 to 318 years (see section 5.1), which did not indicate a difference in the
safety profile of lipegifigrastim in children compared to that in adults. Treatment-related
adverse events were back pain, bome pain and increased neutrophile court (I event each).

averses events were bask pair, usine pair at all interest resumment usual (Levent early). Reporting of systemet adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system.

## 4.9 Overdose

There is no experience with overdose of lipegfilerastim. In the case of overdose, WBC and platelet count should be performed regularly and spleen size should be carefully monitored (e.g. clinical examination, ultrasound).

## 5 PHARMACOLOGICAL PROPERTIES 5.1 Pharmacodynamic properties

## otherapeutic group: Immunostimulants, colony stimulating factors, ATC code:

### Mechanism of action

Mechanism of action Lipediffiguration is a covalent conjugate of fligrastim with a single methoxy polyethylene glycul (PGC) molecule via a carobhylatele linker consisting of glychen, N-acetylneuramist action and N-acetylgalentasamier. The average molecular mass is approximately 30 kB of action and the production and the state of the production of the production of the that regulates the production and release of functional reutrophils from the bone marrow (Figrastin is an un-glycosylated recombinant methors) hy human G-CSF. Lipedifigrastim is a sustained duration form of fligrastim due to decreased renal clearance. Lipedifigrastim blick to human the G-CSF receptor like fligrastim and pelligrastim.

binds to human the C-CS receptor like fligrastim and pegffigrastim. Pharmacodynamic effects Lipsefligrastim and fligrastim induced a marked increase in peripheral blood neutrophil counts within 24 hours, with minor increases in monocytes and/or lymphocytes. These results suggest that the C-CS molety of lipsefligrastim confers the expected activity of this growth factor stimulation of proliferation of haematopoleir progenitor cells, differentiation into mature cells and release into the peripheral blood. This effect includes not only the neutrophil includes but extends to other single lineage and multilineage progenitors and pluripotent haematopoleic stem cells. G-CS also increases the ambitacterial archives of herutophils including the phagecytosis.

<u>Clinical efficacy and safety</u> Once-per-cycle dosing of lipegfilgrastim was investigated in two pivotal randomised, double-blind clinical studies in patients undergoing myelosuppressive chemotherapy. The first pivotal (phase III) clinical study XM22-03 was an active-controlled study in 202 patients with stage II-IV breast cancer receiving up to 4 cycles of chemotherapy consisting of doxorubicin and docetaxel. Patients were randomised 1:1 to receive 6 mg lipegfligrastim or 6 mg pegfligrastim. The study showed non-inferiority of 6 mg ipegfilgrastim to 6 mg pegfilgrastim for the primary endpoint, duration of severe neutropenia (DSN) in the first cycle of chemotherapy (see table 2).

Table 2: DSN, severe neutropenia (SN) and febrile neutropenia (FN) in cycle 1 of study

22-03 (ITT)			
	Pegfilgrastim 6 mg (n = 101)	Lipegfilgrastim 6 mg (n = 101)	
SN.			
ean ± SD (d)	0.9 ± 0.9	0.7 ± 1.0	
LS mean	-0.186		
5 % CI	-0.461 to 0.089		
Į.			
cidence (%)	51.5	43.6	
ł			
cidence (%)	3.0	1.0	

ITT = Intent-to-treat population (all randomised patients)

D = standard deviatio = davs

- confidence interval IS mean (least square mean difference lipegfilgrastim - pegfilgrastim) and CI out

f multivariate Poisson regression analysis he second nivotal (phase III) clinical study XM22-04 was a placeho-controlled study in The second pixetal (phase III) official study MPZ-20 4 was a pixebo-controlled study in 237 patients with non-mail cell lung care receiving up to 4 cycles of chemotherapy may be considered to the considered pixel patients of the considered pixel pixel pixel pixel pixel ligoglifiquation of pixelob. The results of the study are presented in table 3. When the main study was finaled, the incidence of death was 7.2 % [diseated) and 12.5 % (6 mg ligoglifiquation) although after the 360-day follow-up period the overall incidence of death was similar between placeboa and ligoglifiquation (44.8 % and 4.0 % satety population).

## Table 3: DSN. SN and FN in cycle 1 of study XM22-04 (ITT)

	Placebo (n = 125)	Lipegfilgrastim 6 mg (n = 250)		
EN		•		
Incidence (%)	5.6	2.4		
95 % CI		0.121 to 1.260		
p-value		0.1151		
DSN				
Mean ± SD (d)	2.3 ± 2.5	0.6 ± 1.1		
Δ LS mean		-1.661		
95 % CI		-2.089 to -1.232		
p-value		< 0.0001		
SN				
Incidence (%)	59.2	32.1		
Odds ratio		0.325		
95 % CI		0.206 to 0.512		
p-value		< 0.0001		
Δ LS mean (least	square mean difference li	ipegfilgrastim - placebo), Cl and p-va	lue out	

A LS mean (least square mean difference lipegrigrastim - piacebo), Cl and p-val of multivariate Poisson regression analysis

Odds ratio (lipegfilgrastim / placebo), Cl and p-value out of multivariate logistic

gression analysis A nost-authorisation safety study YM22-ONC-40041 was conducted to collect data of

sease progression and mortality in patients with advanced squamous or non-square Ill lung cancer receiving lipegfilgrastim in addition to the platinum-based chemothe creased risk of disease progression or death was not observed with lipegfilgrastim

lysis of anti-drug antibodies of 579 patients and healthy volunteers treated with An analysis of anti-fung patients and health you chusters treated with lingeflignatini, 189 gatients and health you unterest treated with pegflignatini and 121 patients treated with picacho was performed. Drug-specific retailing displants and 121 patients treated with picacho was performed. Drug-specific retailing displants and 121 patients treated with picacho was performed. Drug-specific retailing displants and 121 patients was performed and the picacho was performed and 120 for the subjects receiving pegflignatini and in 1.55 % of the subjects receiving placebo. No neutraling antibodies against liperflignation were observed.

### Paediatric nonulation

r accusance population. The Furnnesh Medicines Agency has deferred the obligation to submit the results of The European Medicines Agency has deferred the obligation to submit the results of studies with Choquies in all subsects of the pediatric population in the treatment of themselves and the properties and prevention of chemothesupy induced leaving continues the properties of the properties and prevention of chemothesupy induced leaving 21 children aged eviewers 2 and 15 years with Every family of tumours or thabdomyosarcoma, lipegifigratism was administrated as a single subcuraneous dose of 100 lug/lig (to to maximum of 6 mg, which is the fixed dose for adults) 24 hours after the properties of the end of the last chemotherapy treatment in week 1 of the regimen. The incidence of FN varied according to age (from 14.3 % to 71.4 %), with the highest frequency in the

oldest age group. The use of three different chemotherany regimens, with varying

myelosuppressive effects and age distributions, complicated the comparison of efficacy

### across age groups (see section 4.2). 5.2 Pharmacokinetic properties

Healthy volunteers In a Studies (XM22-01, XM22-05, XM22-06) in healthy volunteers, the maximum blood concentration was reached after a median of 30 to 36 hours and the average terminal half-life ranged from approximately 32 to 62 hours after a single subcutaneous injection of 6 mg |ipegfligrastim.

injection to this juggingsions.

After subcutaneous injection of 6 mg lipegfligrastim at three different sites (upper arm abdomen and thigh) in healthy volunteers, the bioavailability (peak concentration and area under the curve (AUCI) was 10 word arter subcutaneous injection in the abdomen and in the upper arm. In this limited study WA2CO 6, bioavailability of lipegfligrastim and observed differences among the injection sites were higher in male subjects compared to female subjects.

Nevertheless, pharmacodynamic effects were similar and independent from gender and injection site

## Metabolism

Lipegfilgrastim is metabolised via intra- or extracellular degradation by proteolytic enzymes. Lipegfilgrastim is internalised by neutrophils (non-linear process), then degraded within the cell by endogenous proteolytic enzymes. The linear pathway is likely due to extracellular protein degradation by neutrophil elastace and other plasma

In vitro data indicate that lipegfilgrastim is has little or no direct or immune system mediated effects on CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, and CYP3A4/5 activity. Therefore, lipeefilerastim is not likely to affect metabolism via human cytochrome P450 enzymes

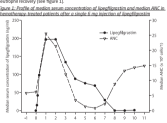
## Special populations

Cancer patients
In 2 studies (XM22-02 and XM22-03) in patients with breast cancer receiving chemotherapy consisting of doxorubicin and docetaxel, mean maximum blood concentrations of 227 and 262 ng/ml were reached after median times to maximu concentrations of 27 and 262 ng/ml were reached after median times to maximum concentration (t<sub>1</sub>) of 44 and 48 hours. The mean terminal half-lives were approximately 2<sup>3</sup> and 31 hours after a single subcutaneous injection of 6 in lipseffiguration draining the first cycle of themotherapy. After a single subcutaneous injection of 6 ing lipseffiguration during the fourth origin. The single subcutaneous injection of 6 ing lipseffiguration during the fourth origin. The single subcutaneous concentrations were lower than observed in the first cycle (inexar values 77 and 111 ng/ml) and were reached after mediant. (a) 6 hours, the mean terminal half-lives in the footh cycle were also proprioritately 39 and 42 hours.

In a study (XM22-04) in patients with non-small cell lung cancer receiving chemotherapy consisting of cisplatin and etoposide, the mean maximum blood concentration of 317 ng/ml was reached after a median  $t_{\rm max}$  of 24 hours and the erminal half-life was approximately 28 hours after a single subcutaneous injection of mg lipegfilgrastim during the first cycle of chemotherapy. After a single subcutaneous injection of 6 me lipeefilerastim during the fourth cycle, the mean maxi concentration of 149 ng/ml was reached after a median t<sub>max</sub> of 8 hours and the mean erminal half-life was approximately 34 hours.

ipegfilgrastim appears to be mainly eliminated by neutrophil-mediated clearance, which becomes saturated at higher doses. Consistent with a self-regulating clearance nechanism, the serum concentration of lipegfilgrastim declines slowly during the hemotherany-induced transient neutronbil nadir and ranidly at the following onset of

eutrophil recovery (see figure 1).



Study days, injection of lipegfilgrastim at day 0 Patients with renal or hepatic impairment
Due to the neutrophil-mediated clearance mechanism, the pharmacokinetics of lipegfilgrastim is not expected to be affected by renal or hepatic impairment

Emitry patients
Limited patient data indicate that the pharmacokinetics of lipegfilgrastim in elderly patients (65 - 74 years) is similar to that in younger patients. No pharmacokinetic data

are available in natients > 75 years

Paediatric population
In a phase 1 study (see section 5.1), using a 10 mg/ml solution for subcutaneous in a place 2 study been sent of 2.1, busing staff integril solution and concentrations (1), were 240 aprilled in the 2 to 6-level group and 10 staff in the 2 to 6-level group, 255 again in the 6 to 12-year group and 224 right in the 2 to 6-level group, 255 again in the 6 to 12-year group after 3 single subcutaneous injection of 100 juggle (maximum 6 long) lipegifigrastin with the first cycle of chemotherapy. The maximum blood concentrations were reached after a median time (t<sub>20</sub>) of 239 flours, 300 hours and 958 flours, respectively, See section 4.5. Overweight patients

Overweight patients
A trend towards a decrease in lipegfilgrastim exposure was observed with increase in weight. This may result in lowered pharmacodynamic responses in heavy patients (9 95 kg), Consequent decrease in efficacy in these patients cannot be excluded on current data.

### 5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, single and repeated dose toxicity and local tolerance. safety pharocology, single and repeated of see toxicity and local loteriance. In a study of toxicity to reproduction and development in rabbits, an increased incidence of post-implantation loss and abortion has been observed at high doses of lipseffjagstalm, filely owing to an exaggeared pharmacolognamic effect specific for rabbits. There is no evidence that lipseffligstalm is teratogenic. These findings are consistent with events thron CGSF and derivatives. Publisher Information on CGSF and derivatives reveal no evidence of adverse effects on fertility and embryo-foetal development in rats or pre-/postnatal effects other than those related to maternal toxicity as well. There is evidence that filerastim and neefilerastim may be transported at low levels over the placenta in rats, although no information is available for

## B DHADMACEUTICAL DADTICULADS

### 6.1 List of excipients

Glacial acetic acid soaium nyarox Sorbitol (E420)

### Water for injections 6.2 Incompatibilities

In the absence of compatibility studies, this medicinal product must not be mixed with other medicinal products

### 6.3 Shelf life

## 6.4 Special precautions for storage

Store in a refrigerator (2°C - 8°C). o not freeze

Keen the pre-filled surings in the outer carton in order to protect from light Lonquex may be removed from the refrigerator and stored below 25°C for a maximum single period of up to 3 days. Once removed from the refrigerator, the medicinal product

### must be used within this period or disposed of. 6.5 Nature and contents of container

Pre-filled syringe (type I glass) with a plunger stopper [poly(ethylene-co-tetrafluoroethylene)-coated bromobutyl rubber] and a fixed injection needle (stainless steel, 29G [0.34 mm] or 27G [0.4 mm] x 0.5 inch [12.7 mm]).

Each pre-filled syringe contains 0.6 ml of solution. Pack sizes of 1 pre-filled syringe with or without safety device (which prevents needle stick injury and re-use).

## Not all pack sizes may be marketed.

6.6 Special precautions for disposal and other handling

The solution should be visually inspected before use. Only clear, colourless solutions without particles should be used

The solution should be allowed to reach a comfortable temperature (15°C - 25°C) for

Vigorous shaking should be avoided. Excessive shaking may aggregate lipegfilgrastim rendering it biologically inactive.

Lonquex does not contain any preservative. In view of the possible risk of microbial contamination, Lonquex syringes are for single use only. Any unused medicinal product or waste material should be disposed of in accordance

### vith local requirement MANUFACTURER

TEVA Pharmaceutical Industries, Ltd 18, Eli Hurvitz St., Industrial Zone Kfar Saba, 4410202, Israel

## ATE OF REVISION OF THE TEXT

