Annex II

Scientific conclusions and grounds for positive opinion presented by the European Medicines Agency
Scientific conclusions

**Overall summary of the scientific evaluation of didanosine and associated names (see Annex I)**

**Background**

Didanosine (2', 3'-dideoxyinosine) is an inhibitor of the in vitro replication of human immunodeficiency virus (HIV) in cultured human cells and cell lines. After didanosine enters the cell, it is enzymatically converted to dideoxyadenosine-triphosphate (ddATP), its active metabolite. In viral nucleic acid replication, incorporation of this 2', 3'-dideoxynucleoside prevents chain extension, and thereby inhibits viral replication. In addition, ddATP inhibits HIV-reverse transcriptase by competing with deoxyadenosine-triphosphate (dATP) for binding to the enzyme's active site, preventing proviral DNA synthesis.

Didanosine and associated names is indicated in combination with other antiretroviral drugs for the treatment of HIV-1 infected patients.

The reference product in EU is Videx EC (200, 250 and 400 mg) hard capsules, first authorised in the United Kingdom (UK) on 19 September 2000.

The application for Didanosine and associated names was considered under Article 10.3 of Directive 2001/83/EC in all Concerned Members States (CMSs).

During the decentralised procedure France and the Netherlands expressed the opinion that bioequivalence had not been demonstrated in the fed conditions as C_{max} was outside the 80-125% acceptability limits1. In addition, the objecting Members States considered that the arguments provided by the applicant did not sufficiently address the consequences of the difference observed in didanosine pharmacokinetics in the fed state between test and reference products.

The decentralised procedure was closed on day 210, with most of the CMSs agreeing with the conclusions of the reference Member State (RMS)’s assessment report except France and the Netherlands which raised a potential serious risk to public health (PSRPH). A referral was thus triggered at the Co-ordination Group for Mutual Recognition and Decentralised Procedures – Human (CMD(h)). The major concern raised by France and the Netherlands could not be solved during the CMD(h) referral and the issue was therefore referred to the Committee for Medicinal Products for Human Use (CHMP).

**Evaluation**

In order to demonstrate the safety and efficacy of Didanosine and associated names in combination with other antiretroviral drugs in the treatment of HIV-1 infected patients, the application dossier was based on two single dose bioequivalence studies, one under fasting state and one under fed state. Both studies were carried out using open label, randomised, two treatment, two sequence, two period, single dose cross-over design. Didanosine and associated names (Gastro-resistant capsules 400 mg) was compared to the reference product Videx EC (Gastro-resistant capsules 400 mg) in 60 healthy adults under fasting conditions. The fed study was conducted during an extended clock-stop.

**Results of the bioequivalence studies**

The primary pharmacokinetic parameters (C_{max} and AUC) were satisfactory in the fasting study with the 90% confidence interval (CI) falling within the standard criteria of 80.00 – 125.00%.
In the fed study, the results were satisfactory in terms of extent of absorption (i.e. AUC) with 90% CI within standard range of 80.00 – 125.00%. However, the 90% CI for C\text{max} was 100.36 – 132.76%. It is acknowledged that these results were outside the standard range of 80 – 125%; however these were within the wider acceptance criteria of 70 – 143% which may be used for highly variable drugs. Of note, bioequivalence in fasted state is considered to be the most important as this product is intended to be taken on an empty stomach.

- **Dosing recommendations for didanosine**

Didanosine is intended to be administrated on an empty stomach as stated in the proposed Summary of Product Characteristics (SmPC): "Didanosine absorption is reduced in the presence of food, and hence Didanosine gastro-resistant capsules should be administrated on an empty stomach (at least 2 hours before or 2 hours after a meal)". Pharmacokinetics studies conducted on Didanosine formulations reveal that administration of the product with food or immediately after food results in decreased in vivo availability of the drug. Since the product is to be administrated at least 2 hours before or after food intake, it is unlikely that the product ingested will be exposed to in vivo conditions, prevailing under fed state.

- **Observed food effect for test and reference products**

As per the current recommendations for modified release formulations\textsuperscript{1,2}, bioequivalence studies in fasted and fed conditions are required. The main purpose of conducting bioequivalence fed study is to exclude food related effects, such as dose-dumping (in particular for gastro-resistant formulations) or failure of protection from acid mediated degradation in the stomach.

The administration of the reference product with a high fat meal significantly decreases the AUC (19%) and C\text{max} (46%) of didanosine. This observation is in line with the results presented in the current application where AUC and C\text{max} of the test product significantly decrease under fed conditions. Therefore, for both the test and the reference product in the fed state the absorption of didanosine decreases which indicate that both products possess similar food effect in terms of reduction in C\text{max} and AUC with no evidence of dose-dumping. The only difference is the magnitude of the decrease, which is less for the test than for the reference product.

The Applicant claimed that the significant decrease observed under fed conditions for C\text{max} for the test product (90% CI outside the standard range) may be attributed to high inter-individual variability with respect to C\text{max} which was 36% in the study conducted under fed conditions. Considering this, the sample size that may be required to meet the standard bio-equivalence criteria would be as high as 232 subjects to obtain power of at least 80%.

The presented data does not give evidence of in vivo dose-dumping from the formulation under fed conditions. Hence, both test and reference products can be considered as having similar food effect in terms of reduction in C\text{max} and AUC.

- **Clinical significance of C\text{max} with Didanosine**

Didanosine has to be first converted intracellularly to its active metabolite ddATP (responsible for antiviral activity) which has a significantly longer intracellular half-life (about 43 hours) as compared with plasma half-life of didanosine. The applicant claimed that differences in plasma concentrations of didanosine are not of clinical relevance as such since it may not result in changes in the intracellular triphosphate concentrations. Initial clinical trials demonstrating efficacy of

\textsuperscript{1} Note for Guidance on the investigation of bioavailability and bioequivalence (EMA/CPMP/EWP/QWP/1401/98)

\textsuperscript{2} Guideline on the pharmacokinetic and clinical evaluation of modified release dosage forms (EMA/CPMP/EWP/280/96/Corr1)
didanosine in treatment of HIV infecting were conducted using buffered tablets\textsuperscript{3,4,5,6,7,8}. The pharmacokinetics data reveals that the plasma concentration ($C_{\text{max}}$) obtained from enteric-coated capsule formulation of didanosine is about 40\% lower as compared to the buffered tablet formulation. This is attributed to the delay in absorption rate of enteric coated formulation which is reflected in the $T_{\text{max}}$ which is about 2 hours for the enteric coated formulation compared to 0.67 hours for the buffered tablet. However, both formulations are equivalent in terms of extent of absorption (i.e. AUC). Therefore, it is considered that the fact that both formulations have been used for the same indications and at similar doses suggests that AUC is more relevant for ensuring efficacy of didanosine in antiviral therapy, and the changes in $C_{\text{max}}$ are unlikely to compromise antiviral efficacy.

The applicant provided literature references which showed that for the action of Didanosine, AUC is the most important parameter\textsuperscript{9,10,11,12,13}. Irrespective of whether Didanosine is taken with or without food, the virological response is based on the total drug exposure. In the current application, the AUC under both fasted and fed conditions was within the acceptance criteria of 80-125\%.

\begin{footnotesize}


\textsuperscript{5} Beltangady M et al. Relation between plasma concentrations of didanosine and markers of antiviral efficacy in adults with AIDS and AIDS related complex. Clinical Infectious Diseases 1993; 16: S26-S31


\textsuperscript{7} Perry CM, Balfour JA. Didanosine: An Update on its Antiviral Activity, Pharmacokinetic Properties and Therapeutic Efficacy in the Management of HIV Disease. Drugs. 1996); 52: 929-962


\textsuperscript{10} Lopez JC et al. A Cohort Study of the Food Effect on Virological Failure and Treatment Discontinuation in Patients on HAART Containing Didanosine Enteric-Coated Capsules (FOODDIe Study). HIV Clin trials. 2006; 7: 155-162


\textsuperscript{12} Berenguer J et al. Didanosine, Lamivudine, and Efavirenz versus Zidovudine, Lamivudine, and Efavirenz for the Initial Treatment of HIV Type 1 Infection: Final Analysis (48 Weeks) of a Prospective, Randomized, Noninferiority Clinical Trial, GESIDA 3903 HIV/AIDS. CID 2008; 47: 1083-1092.

\end{footnotesize}
Grounds for positive opinion

Whereas

- The Committee considered the notification of the referral triggered by the United Kingdom under article 29(4) of Directive 2001/83/EC. the Netherlands and France considered that the granting of the marketing authorisation constitutes a potential serious risk to public health.
- The Committee reviewed all the data submitted by the applicant in order to support the bioequivalence between Didanosine and associated names and the reference product.
- The Committee is of the opinion that bioequivalence has been demonstrated under fasting conditions which is the recommended state for administration of didanosine.
- The Committee noted that in the fed study, both formulations were subject to a food effect reducing plasma concentrations. The bioequivalence studies confirmed that dose dumping did not occur with Didanosine and associated names. The results were satisfactory in terms of extent of absorption (i.e. AUC). The Committee acknowledged that the conventional criterion for bioequivalence for maximum plasma concentration ($C_{max}$) was outside the 80-125% acceptability limits. However, the observed effect of food is lower and the Committee is of the opinion that this is not clinically relevant based on considerations relating to the mechanism of action and, in particular, that Didanosine and associated names is to be administrated on an empty stomach.

the CHMP has recommended the granting of the marketing authorisations for which the summary of product characteristics, labelling and package leaflet remain as per the final versions achieved during the Coordination group procedure as mentioned in Annex III for Didanosine and associated names (see Annex I).