



1 24 September 2015
2 EMA/CHMP/PKWP/36648/2015
3 Committee for Medicinal Products for Human Use (CHMP)

4 **Tacrolimus granules for oral suspension 0.2 and 1 mg**
5 **product-specific bioequivalence guidance**
6 Draft

Draft Agreed by Pharmacokinetics Working Party	April 2015
Adoption by CHMP for release for consultation	24 September 2015
Start of public consultation	1 October 2015
End of consultation (deadline for comments)	1 January 2016

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Comments should be provided using this [template](#). The completed comments form should be sent to PKWPsecretariat@ema.europa.eu.

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Keywords	<i>Bioequivalence, generics, tacrolimus</i>
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10 Tacrolimus granules for oral suspension 0.2 and 1 mg product-specific bioequivalence
11 guidance
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13 Disclaimer:

14 *This guidance should not be understood as being legally enforceable and is without prejudice to the need to ensure that the data submitted in support of a*
15 *marketing authorisation application complies with the appropriate scientific, regulatory and legal requirements.*

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17 Requirements for bioequivalence demonstration (PKWP)*

BCS Classification	BCS Class: <input type="checkbox"/> I <input type="checkbox"/> III <input checked="" type="checkbox"/> Neither of the two Background: tacrolimus may be considered a low solubility compound.
BE Study design <i>in case a BCS biowaiver is not feasible or applied</i>	single dose
	cross-over
	healthy volunteers
	<input checked="" type="checkbox"/> fasting <input type="checkbox"/> fed <input type="checkbox"/> both <input type="checkbox"/> either fasting or fed
	Strength: 1 mg Background: highest strength to be used for a drug with linear pharmacokinetics and low solubility. Higher doses may be needed (multiple 1 mg doses) in case of poor bioanalytical methods.

	Number of studies: one single dose study.
Analyte	<input checked="" type="checkbox"/> parent <input type="checkbox"/> metabolite <input type="checkbox"/> both
	<input type="checkbox"/> plasma/serum <input checked="" type="checkbox"/> blood <input type="checkbox"/> urine
	Enantioselective analytical method: <input type="checkbox"/> yes <input checked="" type="checkbox"/> no
Bioequivalence assessment	Main pharmacokinetic variables: AUC_{0-72h} , C_{max}
	90% confidence interval: 80.00 – 125.00% for C_{max} and 90.00 - 111.11% for AUC_{0-72h} Background: tacrolimus is a narrow therapeutic index drug.

18 * As intra-subject variability of the reference product has not been reviewed to elaborate this product-specific bioequivalence guideline, it is not possible to
19 recommend at this stage the use of a replicate design to demonstrate high intra-subject variability and widen the acceptance range of C_{max} . If high intra-
20 individual variability ($CV_{intra} > 30\%$) is expected, the applicants might follow respective guideline recommendations.