

# MEDICINES CONTROL COUNCIL



DEPARTMENT OF HEALTH  
Republic of South Africa



## ZA CTD and ZA eCTD IMPLEMENTATION

In June 2010 the Medicines Control Council (MCC) announced the intention to implement the South African Common Technical Document (ZA CTD) format which will replace the current MRF1 and any applications still in MBR1 format.

The major milestones of the implementation strategy are as follows:

ZA CTD	Voluntary / recommended	Implementation date
New product applications for registration	1 July 2010	1 June 2011
Post-registration amendments Type A, B and C	1 October 2010	1 June 2011
ZA eCTD	Date	
Pilot phase - new applications for registration		
<i>Step 1: Identify applicants and applications</i>	1 March – 31 May 2013	
<i>Step 2: Run pilot phase</i>	1 <sup>st</sup> June 2013 – 31 March 2015	
Maintenance phase		
<i>Step 3: Start maintenance phase</i>	6 October 2014	
Start Operational Phase		
<i>Step 4: eCTD process open to entire industry for new applications for registration</i>	1 June 2015	
Step 5: All other application types – paper to eCTD	To be advised	
MRF1 format	Last date for acceptance	
New product application for registration	31 May 2011	
Clinical evaluations	31 May 2011	
Post-registration amendments Type A, B and C	31 May 2011	
Post-registration amendments Type A, B and C	<i>Care not acceptable in MBR1 format*</i>	
Conversion of all other MBR1 and MRF1 dossiers	1 June 2016	

\*In exceptional cases, contact the Secretariate for guidance

Pre-registration responses and the submission of post-registration follow-up stability data should comply with the Amendment guideline sections 1 and 6 and may be submitted in the format of the application for registration if still MRF1

**REGISTRAR OF MEDICINES  
MS M HELA**

# Project Plan eCTD implementation at MCC

