



Full wwPDB EM Map Validation Report i

Dec 9, 2020 – 12:41 pm GMT

EMDB ID : EMD-5816
Title : Negative-stain electron microscopy reconstruction of Tetrahymena telomerase (TERT-f, stable conformation)
Authors : , Jiang.J.; , Miracco.EJ.; , Hong.K.; , Eckert.B.; , Chan.H.; , Cash.DD.; , Min.B.; , Zhou.ZH.; , Collins.K.; , Feigon.J.
Deposited on : 2013-12-02
Resolution : 35.00 Å(reported)

This is a Full wwPDB EM Map Validation Report for a publicly released PDB entry.

We welcome your comments at validation@mail.wwpdb.org

A user guide is available at

<https://www.wwpdb.org/validation/2017/EMMapValidationReportHelp>

with specific help available everywhere you see the i symbol.

The following versions of software and data (see [references](#) ①) were used in the production of this report:

EMDB validation analysis : **FAILED**
Validation Pipeline (wwPDB-VP) : 2.13

1 Experimental information i

Property	Value	Source
EM reconstruction method	singleParticle	Depositor
Imposed symmetry	Not Provided	Depositor
Number of images used	156	Depositor
Resolution determination method	OTHER	Depositor
CTF correction method	Each particle	Depositor
Microscope	FEI TECNAI F20	Depositor
Voltage (kV)	200	Depositor
Electron dose ($e^-/\text{\AA}^2$)	40	Depositor
Minimum defocus (nm)	1.2	Depositor
Maximum defocus (nm)	3.7	Depositor
Magnification	68027.0	Depositor
Image detector	TVIPS TEMCAM-F415 (4k x 4k)	Depositor