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Requirementsforaddressingrespiratorymotionincooperative groupclinicaltrials
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Inmid-2006,theNationalCancerInstitute(NCI)publishedupdated guidelinesfor the
useof intensity-modulatedradiotherapy(IMRT)onclinicaltrials,specificallywhen
thetargetvolumeincludes thethoracicregionorotherareasinwhichrespiratorymotion
couldhaveasignificanteffect.Inadditiontorequiringheterogeneitycorrections, the
NCInowrequires thattheclinicalprotocols address the localization and immobilization of
boththepatientandthetumor. Imaging must be performed in a manner that provides a
representation of the target volume without motion artifact. Procedures must be defined
to document reproducible daily position of the patient and target. Some form of
credentialing is required.

TheRadiologicalPhysicsCenter(RPC)hasbeenenlistedtoparticipatein the
credentialingprocessforinstitutions participatingincertaincooperative grouptrialsin
whichrespiratorymotionis an issue. To accomplish this, theRPChas constructed
severalphantomsthatmimicthethoracicandabdominalregions, and which can be placed
onamovingplatformtosimulate respiratorymotion.The combination of phantom and
movingplatformhave been used to evaluate compensation techniques for respiratory
motionatseveralinstitutions. Thesetechniquesemployedandtheresultsofthese
measurementswillbedescribedaswellas those reported in the literature.

Learningobjectives:

1. Review the structure supporting cooperative group clinical trials in the US.
2. Become familiar with the NCI guidelines for the use of IMRT in clinical trials.
3. Learn about the effects of respiratory motion during thoracic treatments.
4. Understand the information derived from the RPC's moving anthropomorphic
phantoms.

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