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Requirementsforaddressingrespiratorymotionincooperative groupclinicaltrials
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Inmid-2006,theNationalCancerInstitute(NCI)publishedupdated guidelinesfor the useof intensity-modulatedradiationtherapy(IMRT)oncologicaltrials,specificallywhen thetargetvolumeislocatedinthoracicregionorotherareasinwhichrespiratorymotion couldhaveasignificanteffect.Inadditiontorequiringheterogeneitycorrections, the NCInowrequires thattheclinicalprotocolsaddresshelicalizationandimmobilizationof boththepatientandthetumor.Imagingmustbe performedinamannerthat providesa representationofthetargetvolume withoutmotionartifact.Proceduresmustbe defined todocumentreproducibledailypositionofthepatientandtarget.Someformof credentialingisrequired.

TheRadiologicalPhysicsCenter(RPC)hasbeenenlistedtoparticipateinthecredentialingprocessforinstitutions participatingincertaincooperative grouptrialsin whichrespiratorymotionis anissue.Toaccomplishthis, theRPChas constructed severalphantomsthatmimicthoracicandabdominalregions,which canbeplaced onamovingplatformtosimulate respiratorymotion.The combination of phantomand movingplatformhave be usedto evaluatecompensationtechniquesforrespiratory motionatseveralinstitutions.Thesetechniquesemployedandtheresultsofthese measurementswillbedescribedaswellasthose reportedintheliterature.

Learningobjectives:

1. Reviewthestructure supportingcooperativegroupclinicaltrials inthe US.
- 2.BecomefamiliarwiththeNCIguidelinesfortheuseofIMRTinclinicaltrials.
3. Learn about the effectsof respiratorymotionduring thoracic treatments.
- 4.Understand theinformationderivedfrom theRPC's movinganthropomorphic phantoms.

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