TomD.Humphreys,TinaM.Jayaweera,RobertL.Lee LawrenceLivermoreNationalLaboratory,Livermore,California

1.INTRODUCTION

Manyhazardousatmosphericreleases involvechemicalreactionsthatoccurwithinafew kilometersofthesource.Reactionswith commonlyoccurringatmosphericcompounds suchastheOHradical,cantransformand potentiallyneutralizeoriginalreleasecompounds. Especiallyinthesecases,a ccuratelyresolving flowaroundnearbystructuresandover surroundingtopographycanbecriticalto correctlypredictingmaterialdispersion, and thus, theextentofanyhazard.

2.PROGRAMCOMPONENTS

Thefiniteelementcomputationalfluid dynamics(CFD)codeFEM3MP(Chanand Stevens,2000)runswithahorizontaldomain sizeontheorderofkilometers.amaximumgrid resolutionontheorderofonemeter, and a time stepontheorderofonesecond.Individual buildingswithinthedomainareresolved slipboundaryconditions.Windswitharound0.5 -5m/secatbuildinglevelhavetransittimes acrossthecomputationaldomainaroundanhour ortwo. The codes olves for a steady or time dependent/velocityfield,dependingontheuseof aRANSo rLESturbulencemodel. These velocityfieldsadvectpassivetracersas continuousscalars.(Humphreys, etal., 2003)

Totrackchemicalreactionsamongthese passivetracers.weincorporatedintothecode thechemicalreactionsolverSMVGearII (Jacobson, 1999). This solver is called a tevery timestepinalternationwiththevelocitysolver, evolvingthechemicalcompositionateachgrid pointwithnoreferencetoneighboringpoints.

3.PHYSICALCONSIDERATIONS

Chemicalreactionsrelevanttoflow swithin thiscomputationalsystemmustthusoccuron timescalesbetweenonesecondandacouple hours.Reactionsoccurringfasterthanone secondarecombinedwithslowerreactions.while reactionstakinglongerthanacouplehoursare omitted.Reaction swithonlyminorproduct pathwaysarealsoomitted.Whilephotolytic decayiseasilyincludedinthesimulation,rates are highly dependent on temporal variations, e.g.,

cloudcover, and may need to be modified for eachsituation.

Nochangeshavebeenm adeatpresentto linkthereactionsandtheenergyequation, i.e., reactionsneitherabsorbheatfromnorinjectheat intotheflow. This is are a sonable assumption for thediffuse, non -energetic reaction stypical of the atmosphere.

4.ASSEMBLYAND **TESTING**

Asetofreactionsandreactionconstantsina textfileisautomaticallyprocessedtoproduce Fortranprogramfilesencodingthechemical mechanism. This processing is done by scripts writtenbyPeterConnellatLLNL(Dickinson, 1976).(By"mec hanism", Irefertoasetof interconnectedreactionsandreactionratesfor them.)Theseareincludedinthecodeduring compilation. This fixes the mechanism within the executableprogram, but means that new mechanismscanbeimplementedwithno changestotherestofthecode. Mechanisms generatedsofarmodelthedecayofcompounds relevanttotheproductionorreleaseofchemical weapons, but application stoindustrial compoundsorairqualityissuesarealsobeing investigated.

Usingatrialmecha nismforthedecayof dimethylether(basedonGood,2000),theSMV Gearllinterfacesubroutineproducedthesame resultsasaseparate"box"(one -dimensional) reactionmodel. These results were again duplicatedwithinthejointFEM3MP/SMVGearII codeinadomaininwhichallcellswereidentical andtherewasnovelocityofdiffusion. Given the constraintsonthemechanismandthevalidation oftheCFDandchemistrysolversseparately,we conclude that the joint code is accurate and valid. Anyrefer encestodataofthechemicaland spatialevolutionofanatmosphericreleasewould bemostwelcome.

5.DEMONSTRATIONS

Simulationsofreactingplumeswithdifferent chemicalmechanismsrevealfeaturesrelevantto hazardassessmentinanurbanenviron The ground of the image in Figure 1 is colored by the(logofthe)concentrationoftheinitialrelease material. Note the entrainment of the plume in

thewakeofthetowerinfrontofthesource. The 3Dgrayisoconcentration represents one of the reaction products. The extent of its maximum concentration are ais much less than that of the main plumed ue to a moderate reaction rate and the spread of the main plumed ue to obstacle sin the flow. Faster reactions could significantly limit the spread of the main plume.

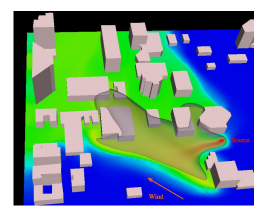


Fig.1 Inthisreactingplume, the ground is colored by the concentration of the initial release material. The gray is osurface is that of a reaction product.

Plumesofreactionproductshavefeatures notsharedbythemainplume. Asseenin Figure2,themaximumoftheproductplumecan beseparatedfromthesource.

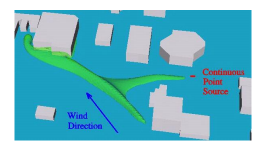


Fig.2 Thisisosurfaceofabreakdownproductis displacedfromthemainplumesource.

Inaddition, product maximum scanbe noncontiguous, tending to formin egions of slower-moving air. At the center of Figure 3, we see a product maximum in the wake to the side of a building.

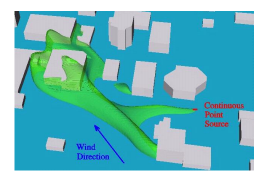


Fig.3 Anisolatedlocalmaximumofreactionproduct concentrationformsintherecirculationregionon thesideofabuilding.

Solvingamechanismwiththirteenspecies andtenreactionsincreasedthecomputational costsofatimestepbyaboutonethirdover advectionanddiffusionofthirteenspecies withoutreactions. Weestimatethatsolvingfora velocityfieldandchemicall yreactingspecies dispersedwithinitwouldtakeabouttwiceas muchtimeassolvingforthevelocityfieldalone.

6.CONCLUSIONS

Accuratepredictionofmaterialdispersion aroundcomplexgeometriesnearthesourceof anatmosphericreleaserequires high-resolution computation. Further complications arise if the compounds released under gochemical reactions which could alter the extent of the main plume. The reaction products form dispersion patterns separate from, and of tenmore complicated than, the original plume.

Directionsforfutureworkincludeexpanding thelibraryofchemicalreactionmechanisms, addingcapabilitiesforaqueousand heterogeneousreactions, and integrating this model within larger -scale models. We plan that the larger -scale models will provide meteorological and chemical boundary conditions, and that this model could provide a source termin larger -scale models, both for momentum and for dispersed compounds.

7.REFERENCES

Chan, S, and D. Stevens, 2000: Validation of two CFDUrban Dispersion Models Using High Resolution Wind Tunnel Data. Third Int. Sym. on Environ. Hydraulics, Arizona State University, Tempe, AZ, Dec 5-8, 2001, 107.

Dickinson,R.P.,R.J.Gelinas,andR.L.Tarp, 1976:AGeneralKineticsPreprocessor Program.LawrenceLivermoreNational LaboratoryPreprint,UCRL78809. Good,D.A.,andJ.S.Francisco,2000: TroposphericOxidationMechanismof DimethylEtherandMethylFormate.J.Phys. Chem.A,104(6),1171 -1185.

Humphreys,T.,S.Chan,andR.Lee, 2003: ValidationofCFDNear -buildingDispersion withUrban2000DataforSteadyandTime dependentBoundaryConditions.Seventh AnnualGMUTransportandDispersion ModelingWorkshop,Fairfax,VA,June17 -19,2003.

Jacobson, M.Z., 1999: Improvement of SMVGearII on Vector and Scalar Machines Through Absolute Error Tolerance Control. *Atmos. Environ.*, 32,791 -796.

ACKNOWLEDGEMENTS

WethankPhilipCameron -SmithandPeter ConnellfortheirinvaluablehelpwiththeSMV GearIIsubroutines,themechanismge neration scripts,andtheconstructionofaninitialtrial mechanism.Thisworkwasperformedunderthe auspicesoftheU.S.DepartmentofEnergyby theLawrenceLivermoreNationalLaboratory undercontractNo.W -7405-Eng-48.