

Status of MPEXS-DNA

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Background

- Geant4-DNA simulation needs a long computation time
 - Tracks a large number of particles and molecular species with the Monte Carlo method
- MPEXS-DNA: A nanodosimetry simulator running on GPU
 - Succeeded to boost up computing performance for physics and chemistry simulation drastically
 - Can accelerate simulation studies on nanodosimetry



- Next step:
 - Implementing an alternative model for chemistry simulation
 - Quantitative DNA damage estimation

MPEXS-DNA: A Nanodosimetry simulation on GPU

- Rewrite Geant4-DNA (10.2.p3) in CUDA
 - EM Physics with lower energy range (down to meV)
 - Calculate local energy loss and generate primary molecules like H_2O^* and $\text{H}_2\text{O}^{+/-}$
 - Radiolysis of water
 - Diffusion and reactions for molecular species
 - Adopting the *step-by-step* method
 - The biological stage not yet implemented

- Collaboration between  and 

- Published a paper on MPEXS-DNA from Medical Physics

MEDICAL PHYSICS

The International Journal of Medical Physics Research and Practice

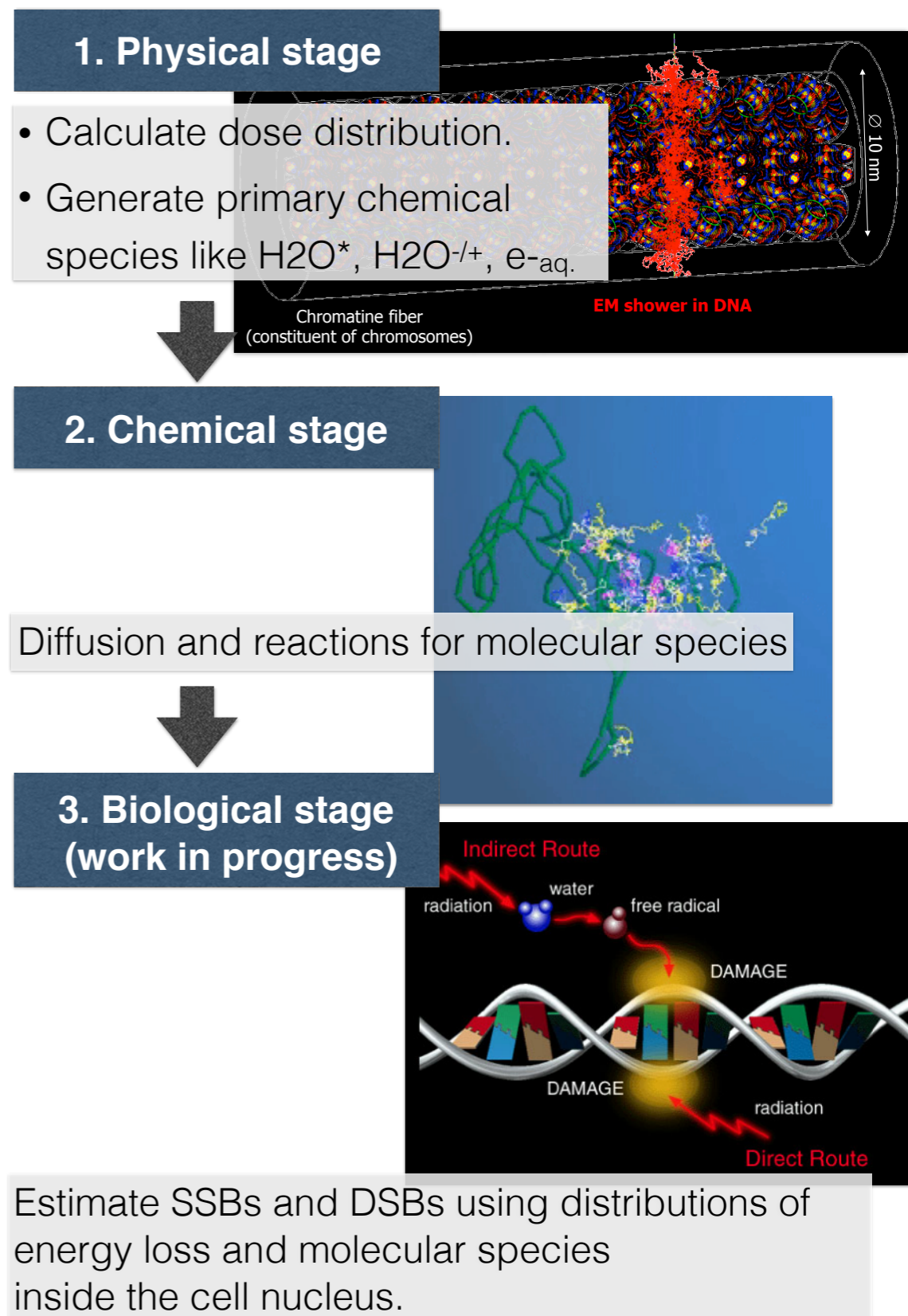
Research Article

MPEXS-DNA, a new GPU-based Monte Carlo simulator for track structures and radiation chemistry at subcellular scale

Shogo Okada , Koichi Murakami, Sebastien Incerti, Katsuya Amako, Takashi Sasaki

First published: 28 December 2018 | <https://doi.org/10.1002/mp.13370>

doi:10.1002/mp.13370



Physical Stage

- Most of the physics lists for Geant4-DNA are covered
- Some physics models released after 10.2.3 are not implemented
 - The CPA100 model (elastic scattering/ionization/excitation)
 - Physics interactions for DNA components
 - ... → will be implemented soon

✓: Available, ✗: Later

Physics lists for G4DNA (10.5.1)	MPEXS-DNA
G4EmDNAPhysics	✓
G4EmDNAPhysics_option1	✓
G4EmDNAPhysics_option2	✓
G4EmDNAPhysics_option3	✓
G4EmDNAPhysics_option4	✓
G4EmDNAPhysics_option5	✓
G4EmDNAPhysics_option6	✗
G4EmDNAPhysics_option7	✓
G4EmDNAPhysics_option8	✗

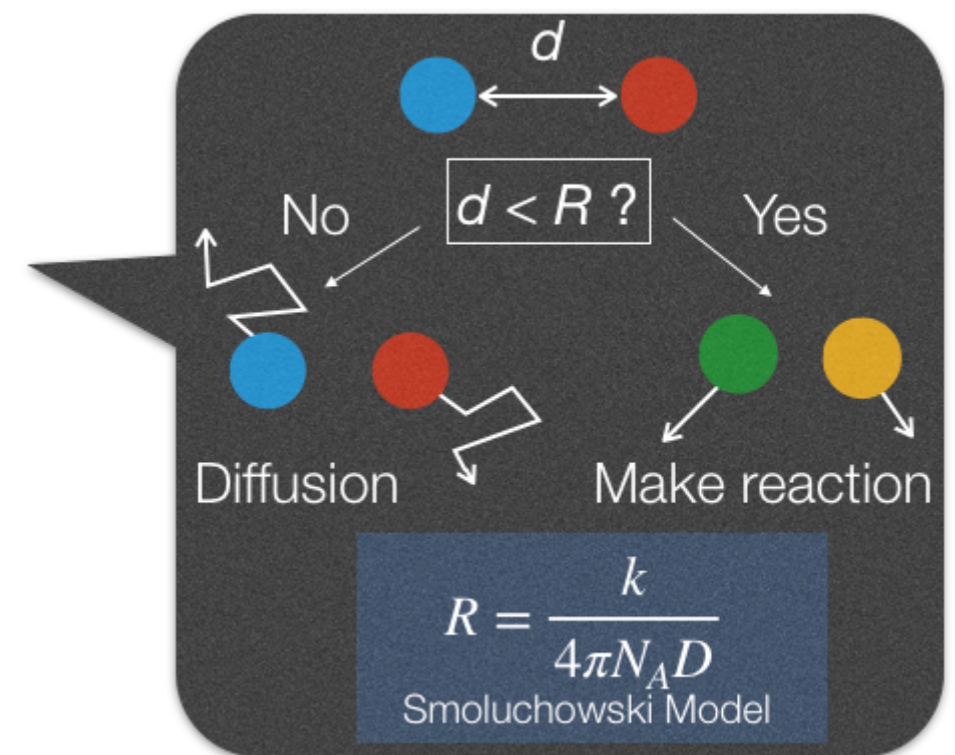
Physico-Chemical Stage

- All processes in Geant4-DNA are implemented in MPEXS-DNA:
 - Dissociation, relaxation, and electron-hole recombination for $\text{H}_2\text{O}^+/\text{H}_2\text{O}^-$, H_2O^* molecules
 - Thermalization for electrons

Electronic state	Process	Dissociation channel	Fraction (%)
Ionization state	Dissociative decay	$\text{H}_3\text{O}^+ + \bullet\text{OH}$	100
Excitation state: A1B1	Dissociative decay	$\bullet\text{OH} + \text{H}\bullet$	65
	Relaxation	$\text{H}_2\text{O} + \Delta\text{E}$	35
Excitation state: B1A1	Auto-ionization	$\text{H}_3\text{O}^+ + \bullet\text{OH} + \text{e}^-_{\text{aq}}$	55
	Dissociative decay	$\bullet\text{OH} + \bullet\text{OH} + \text{H}_2$	15
	Relaxation	$\text{H}_2\text{O} + \Delta\text{E}$	30
Excitation state: Rydberg, diffusion bands	Auto-ionization	$\text{H}_3\text{O}^+ + \bullet\text{OH} + \text{e}^-_{\text{aq}}$	50
	Relaxation	$\text{H}_2\text{O} + \Delta\text{E}$	50
Dissociative attachment	Dissociative decay	$\bullet\text{OH} + \text{OH}^- + \text{H}_2$	100
Electron Hole Recombination	Dissociative decay	$\bullet\text{OH} + \bullet\text{OH} + \text{H}_2$	15
	Dissociative decay	$\bullet\text{OH} + \text{H}\bullet$	55
	Relaxation	$\text{H}_2\text{O} + \Delta\text{E}$	30

Chemical Stage

- Simulate diffusion and chemical reactions for molecular species from 1 ps to 1 us after irradiation
 - Reactions based on the Smoluchowski theory
 - Molecular diffusion with a **step-by-step** method
 - Dynamical time-stepping
 - Calculate encounter probability using the Brownian-bridge method
- Two sets of parameters:
 - G4EmDNACChemistry and G4EmDNACChemistry_option1



Diffusion Coefficients [$\times 10^{-9}$ m²/s]

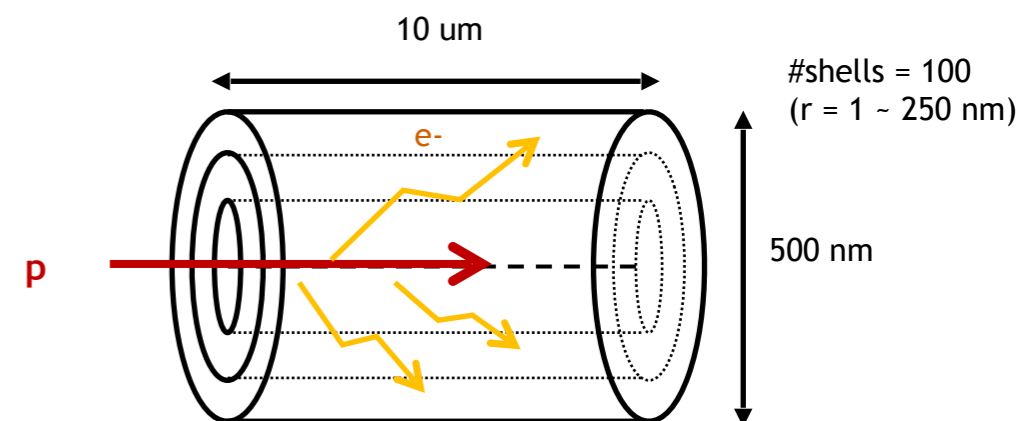
	G4EmDNACChemistry	G4EmDNACChemistry_opt1
H ₃ O ⁺	9.0	9.46
H•	7.0	4.8
OH ⁻	5.0	5.3
e ⁻ _{aq}	4.9	4.9
H ₂	4.8	4.8
•OH	2.8	2.2
H ₂ O ₂	2.3	2.3

Reaction Rate Constants [$\times 10^{10}/(\text{M}\cdot\text{sec})$]

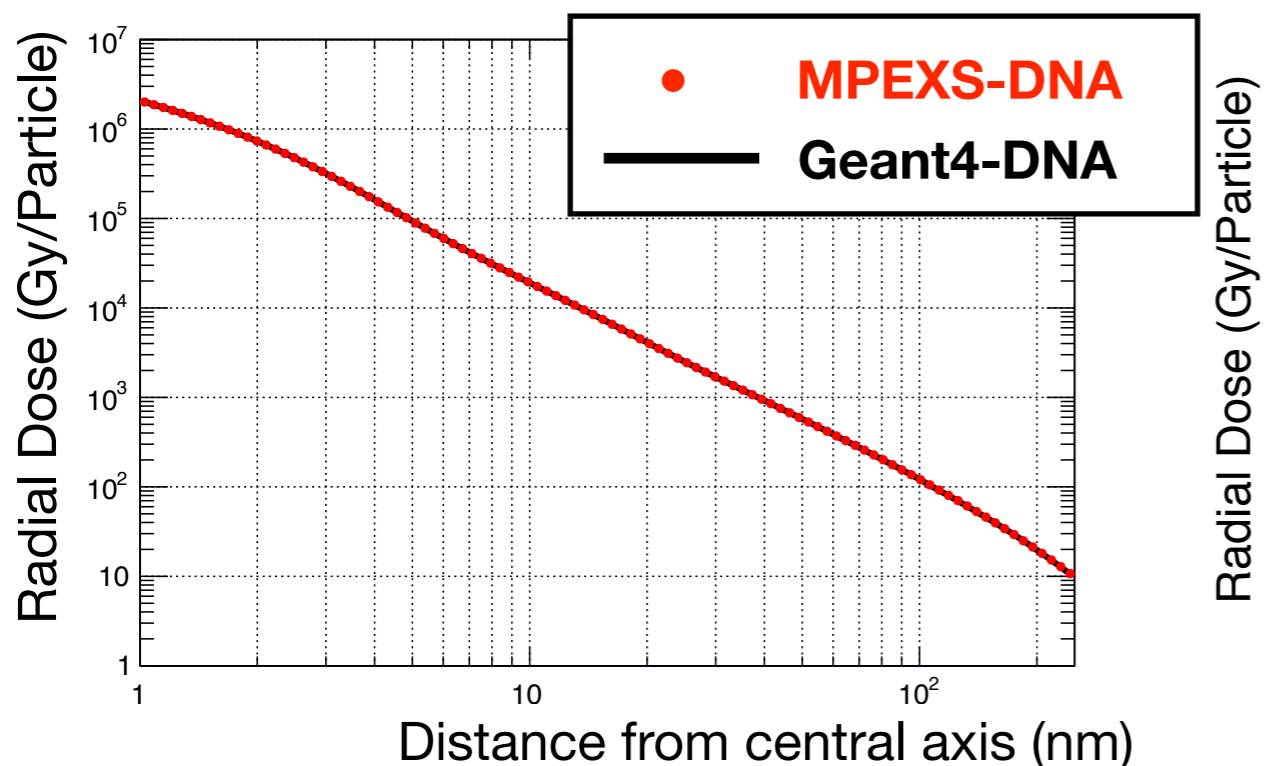
	G4EmDNACChemistry	G4EmDNACChemistry_opt1
2e ⁻ _{aq} + 2H ₂ O → H ₂ + 2OH ⁻	0.50	0.636
e ⁻ _{aq} + •OH → OH ⁻	2.95	2.95
e ⁻ _{aq} + H• + H ₂ O → OH ⁻ + H ₂	2.65	2.50
e ⁻ _{aq} + H ₃ O ⁺ → H• + H ₂ O	2.11	2.11
e ⁻ _{aq} + H ₂ O ₂ → OH ⁻ + •OH	1.44	1.10
•OH + •OH → H ₂ O	0.44	0.55
•OH + H• → H ₂ O	1.44	1.55
H• + H• → H ₂	1.20	0.503
H ₃ O ⁺ + OH ⁻ → 2H ₂ O	1.43	11.3

Physics Verification/Validation

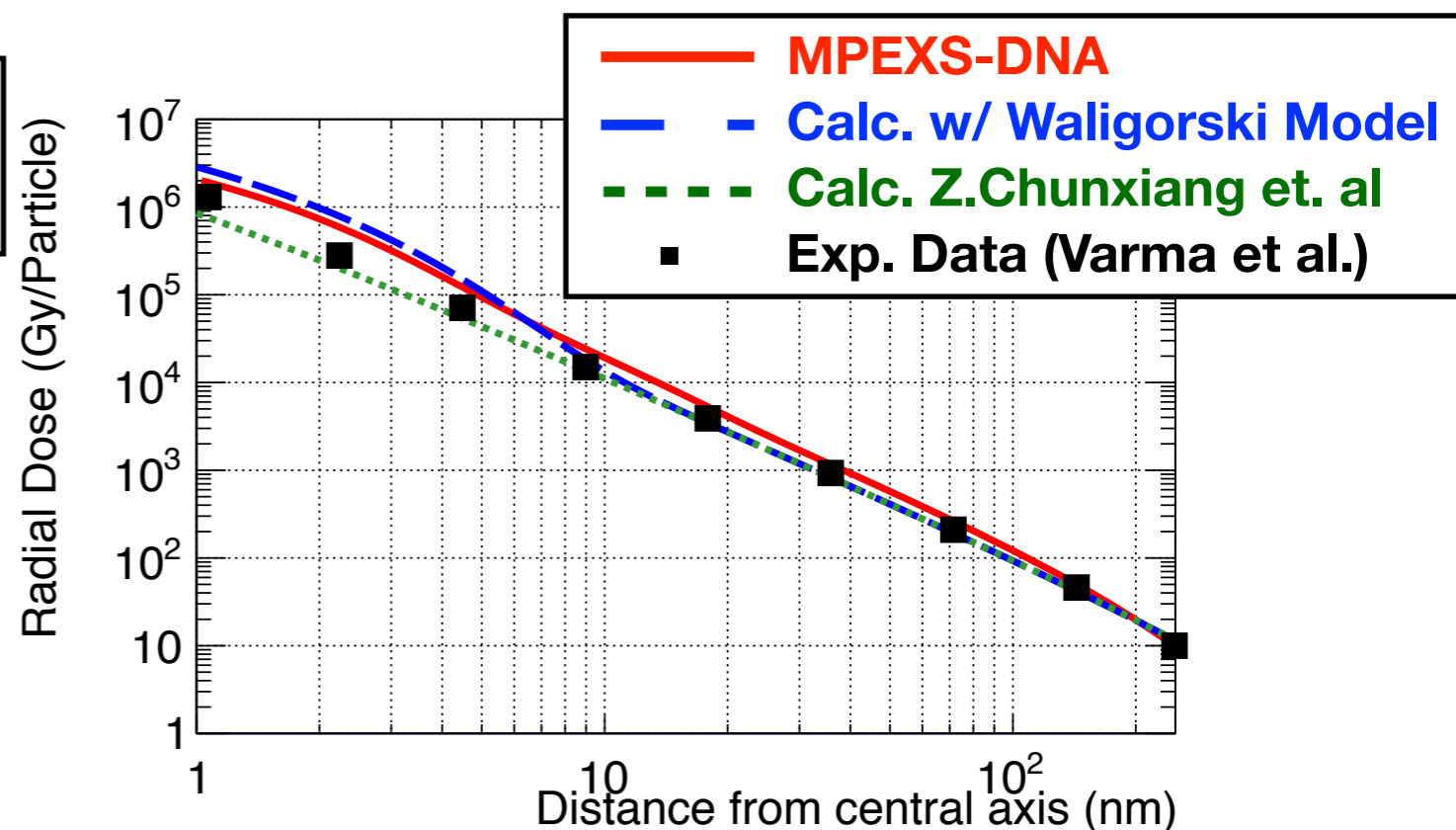
- Compared radial dose distribution of MPEXS-DNA to Geant4-DNA and experimental data
 - Physics List: G4EmDNAPhysics_option4



A: Radial dose distribution:
MPEXS-DNA vs Geant4-DNA
 (2.57 MeV/u Oxygen atoms)



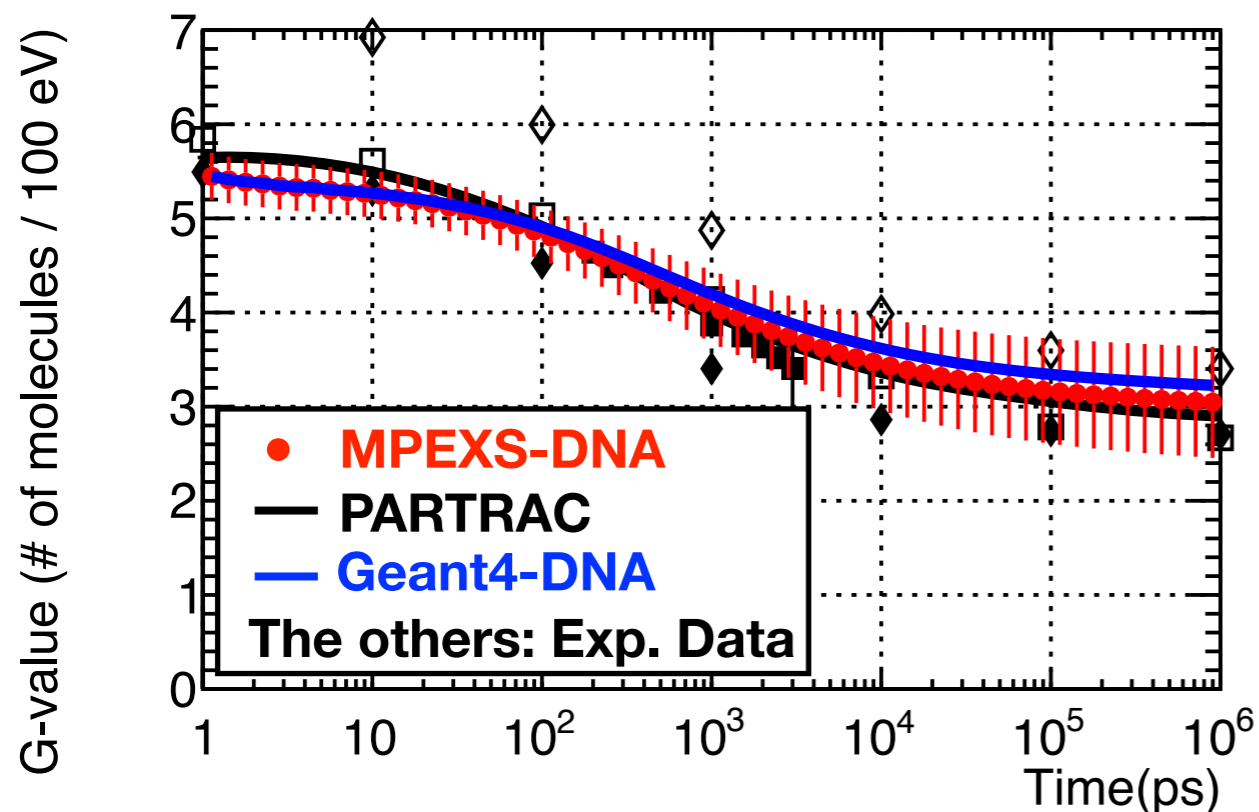
B: Radial dose distribution:
MPEXS-DNA vs measurement data
 (2.57 MeV/u Oxygen atoms)



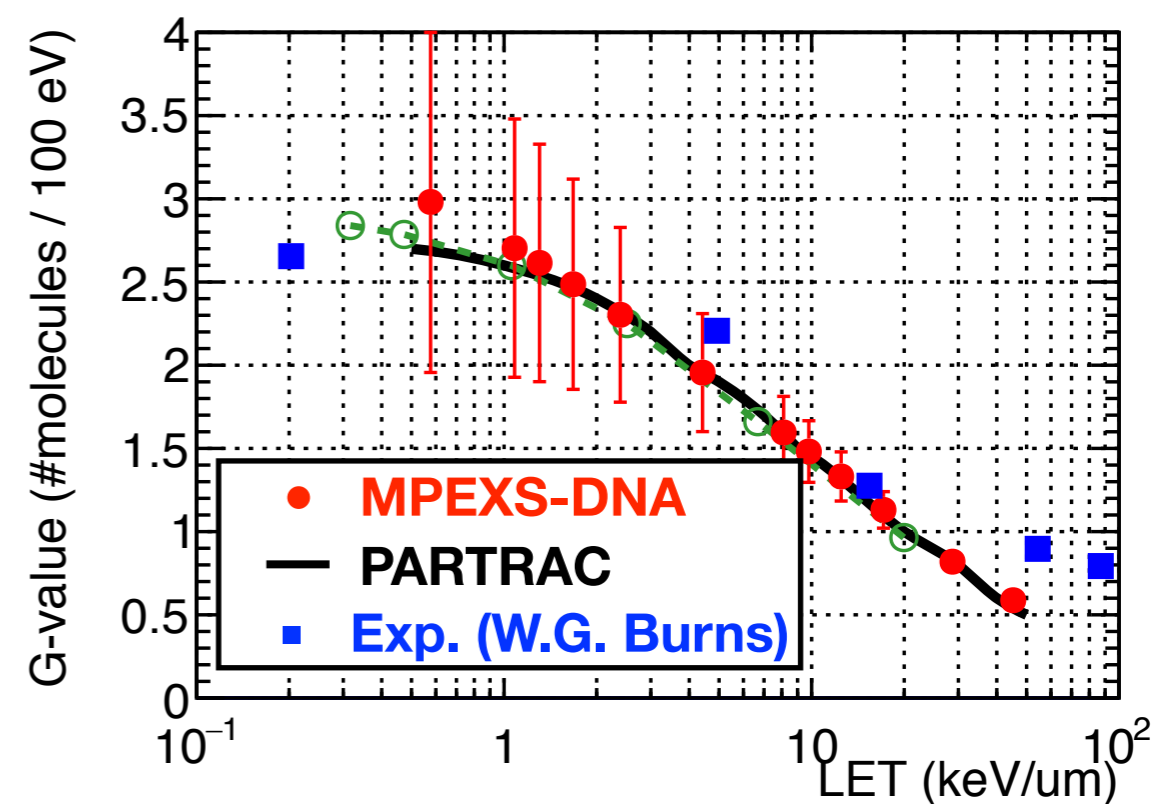
Chemistry Verification/Validation

- Compared G-value plots of MPEXS-DNA to the other MC simulations and measurement data
 - Physics List: G4EmDNAPhysics_option4 and G4EmDNAChemistry

A: G-value time profile for $\cdot\text{OH}$ radicals from 1 ps to 1 μs after irradiation of e^-



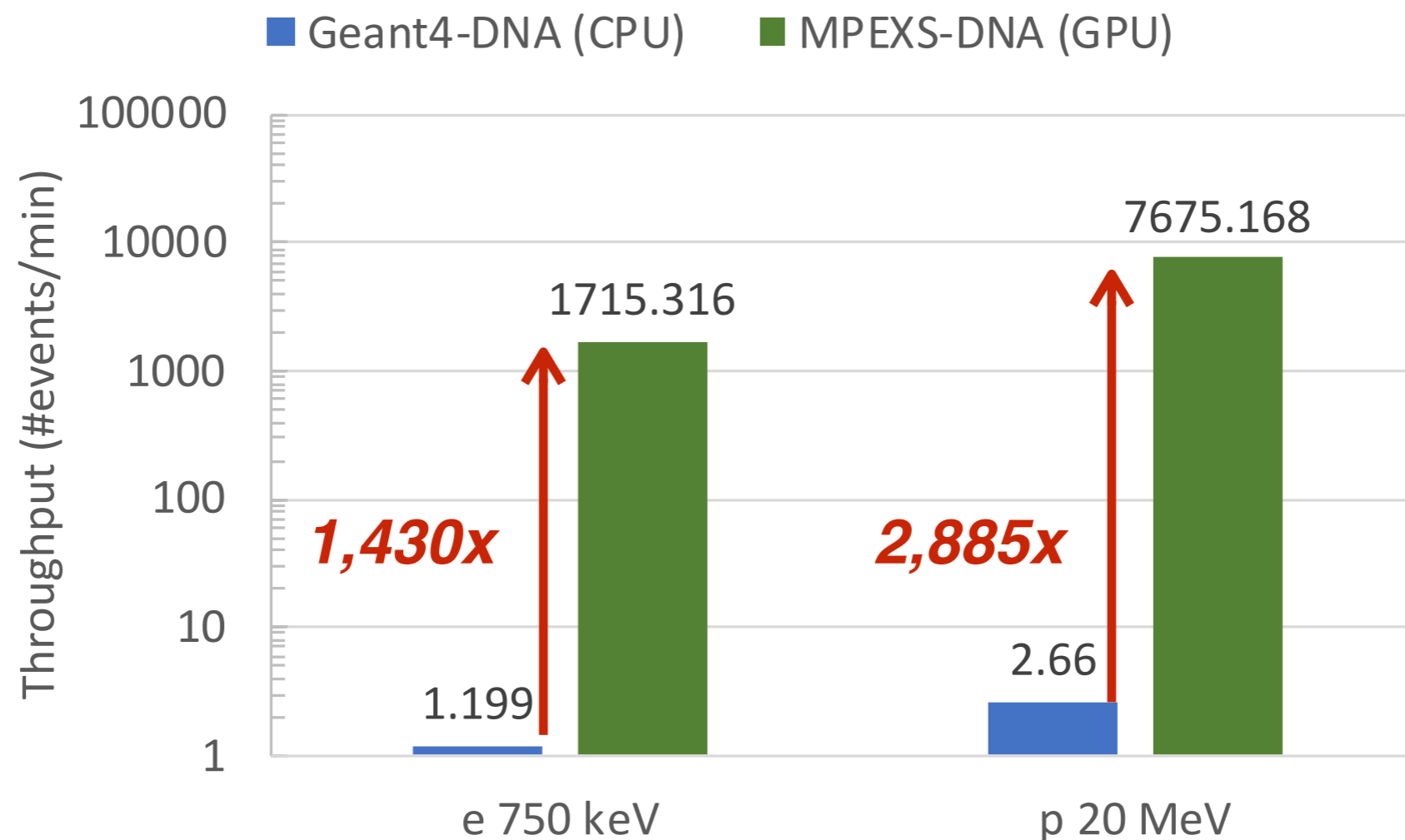
B: LET dependency of G-value for $\cdot\text{OH}$ radicals at 1 μs after proton irradiation



MPEXS-DNA Performance

- Up to **2,900 times** faster than Geant4-DNA with single-core of CPU
 - ~ 4 days (CPU) -> ~ **2 min. (GPU)**
 - **1 GPU unit has the equivalent computing power as 2,900 CPU cores**

Comparisons of event number processed per 1 min.



- GPU: NVIDIA, TITAN V
- CPU: Intel, Xeon E5-2643 v2, 3.5 GHz

MPEXS-DNA Functionalities

	Status
Physics Stage	<ul style="list-style-type: none">• Most of physics models in G4DNA can be run<ul style="list-style-type: none">• The CPA100 model will be implemented soon
Physico-chemical Stage	<ul style="list-style-type: none">• Implemented all processes in G4DNA:<ul style="list-style-type: none">• Dissociation, Relaxation, and Electron hole recombination for H_2O^*, $\text{H}_2\text{O}^+/\text{H}_2\text{O}^-$• Thermalization for electrons
Chemical Stage	<ul style="list-style-type: none">• Simulate diffusion and reactions with the same manner as G4DNA<ul style="list-style-type: none">• Two-parameter sets: G4EmChemistry and G4EmChemistry_option1

**Recent activities
on Geant4-DNA and MPEXS-DNA**

Implementation of An Alternative Model for Chemistry Stage

- **Plante's approach**
 - Ref.) I. Plante, DOI: 10.1007/s00411-011-0367-8
 - Based on the Smoluchowski theory for reactions
 - Handling electrostatic interactions and spin effects
 - Computing encounter probability by solving the diffusion equation with Green's function
 - Simulating reactions with homogeneously high-concentrated species
 - ex. radical scavengers
- **Implementation of Plants's approach in MPEXS-DNA**
 - Diffuse molecules *step-by-step* by Brownian motion
 - Adopting static time-stepping
 - **Implemented using G4DNA (10.5 and 10.6 beta) tentatively**
- **Plante's approach with *IRT* for Geant4-DNA**
 - Developing by Wook (CENBG) and Hoang (IRNS) with the support from Jose (UCSF)

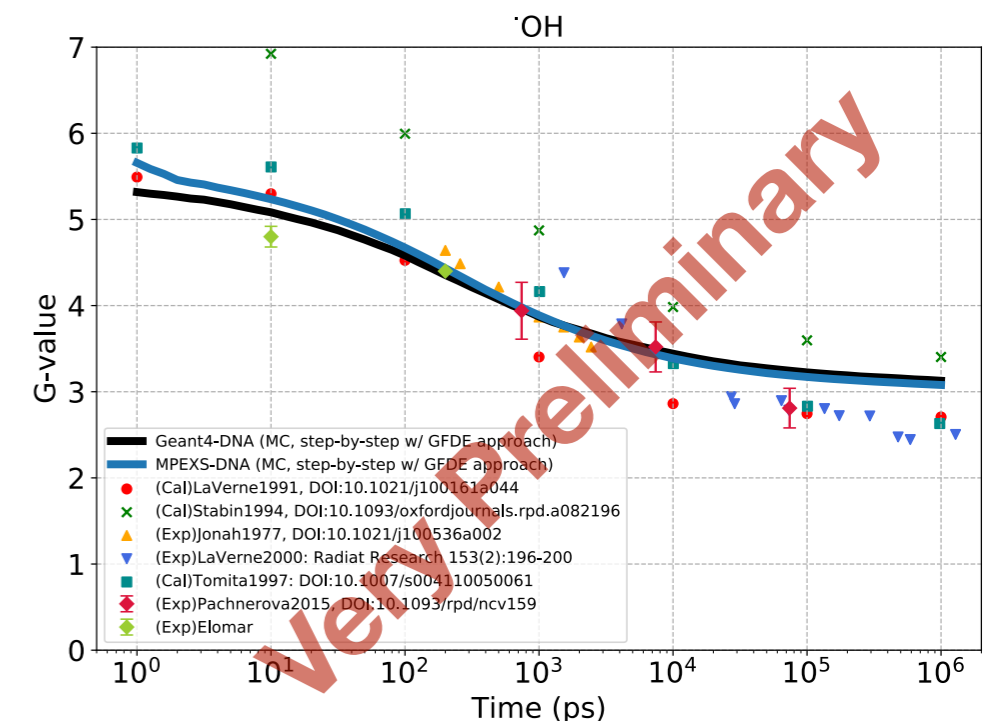
Radiat Environ Biophys (2011) 50:389–403
DOI 10.1007/s00411-011-0367-8

ORIGINAL PAPER

A Monte-Carlo step-by-step simulation code of the non-homogeneous chemistry of the radiolysis of water and aqueous solutions. Part I: theoretical framework and implementation

Ianik Plante

G-value time profile for $\cdot\text{OH}$ radicals from 1 ps to 1 μs after irradiation of 750 keV electrons



Estimation of DNA Damages

- Geant4-DNA calculates early radiation damages using a complex geometry of the cell nucleus
 - Process flow:
 1. Calculate distributions of energy loss and molecular species
 2. Score the positions at which physical and chemical interactions occurred with DNA molecules in a cellular geometry
 3. Evaluate DNA damages with higher geometrical accuracy
 - Reproduce experimental data and the other simulation results well
- We are trying to estimate DNA damages with an alternative approach
 - Simple cell model for geometry
 - Clustering model for DNA damages

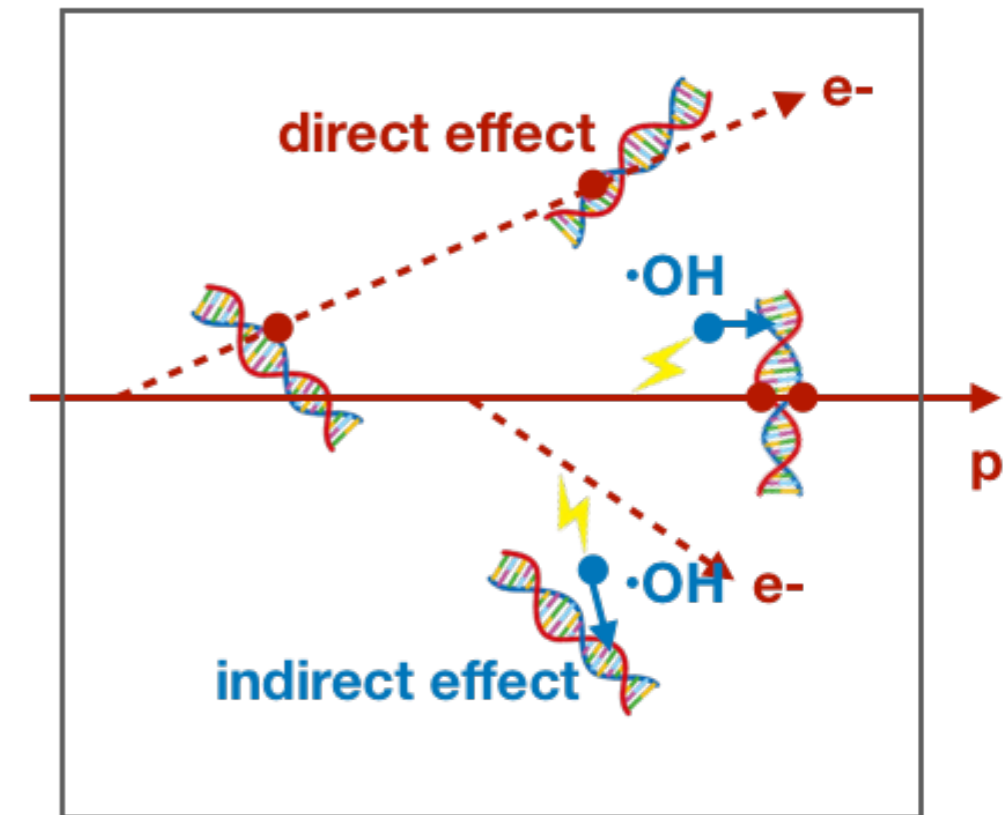
Quantitative Estimation of DNA Damage Using Clustering Model

- “clustering”: An extended example for Geant4-DNA
 - Evaluates DNA damage induced by direct effects
 - Ref.) DOI:10.1016/j.cmpb.2010.12.012

- We extend the “clustering” application

- Estimate DNA damages by **indirect effects** as well as direct effects

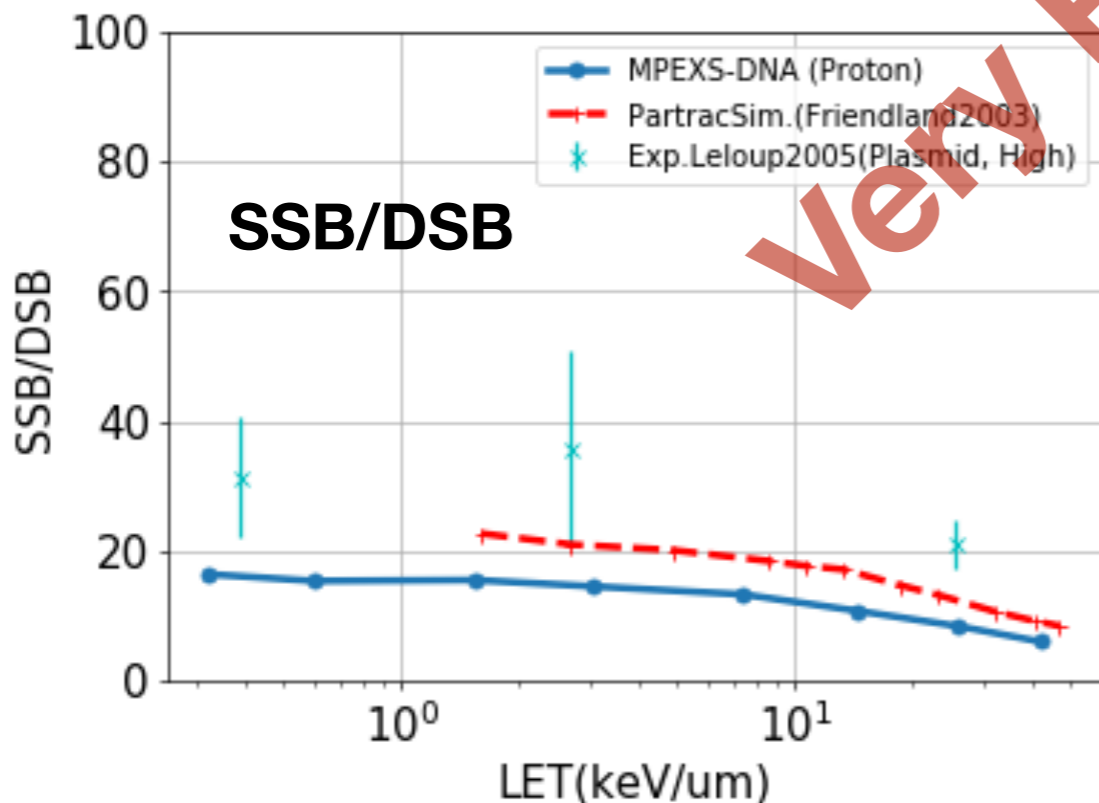
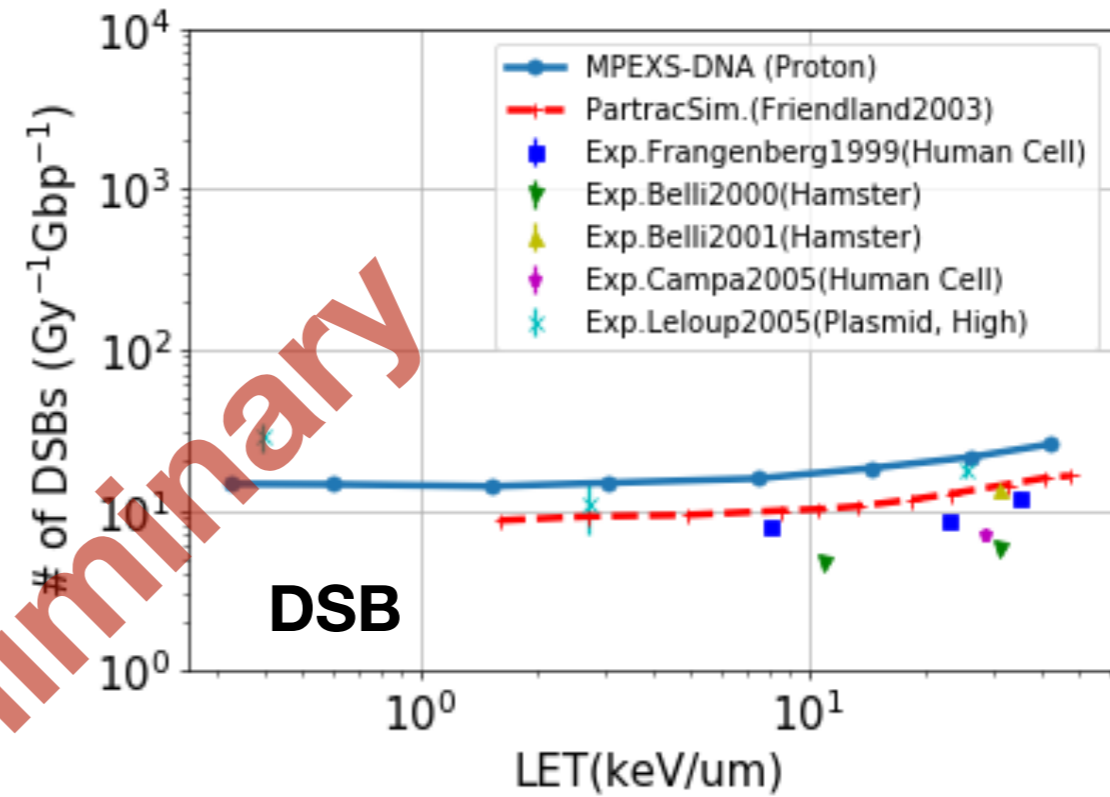
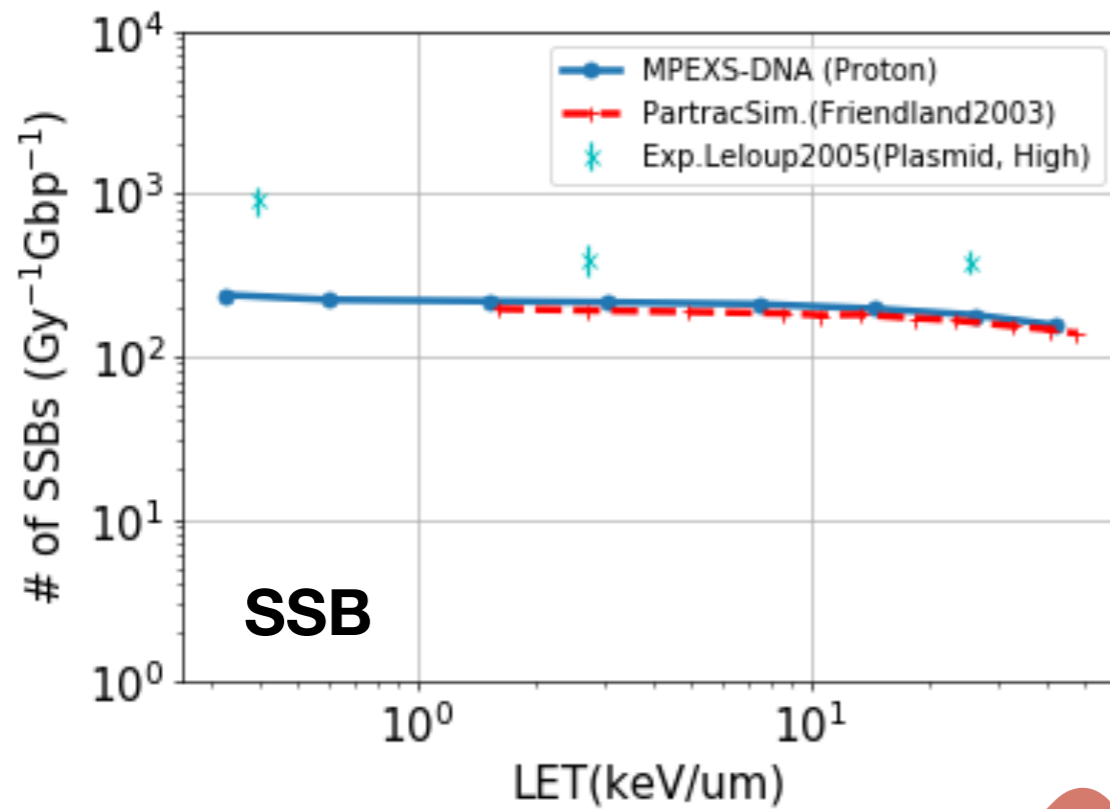
- Assume that DNA molecules are homogeneously diluted in a box geometry (Simple cell model)



- Simulate track structure of charged particles and molecular species
 - Using the Plante’s approach for chemistry simulation

- Sample the positions at which physical and chemical reactions with DNA components occur, then evaluate DNA damages

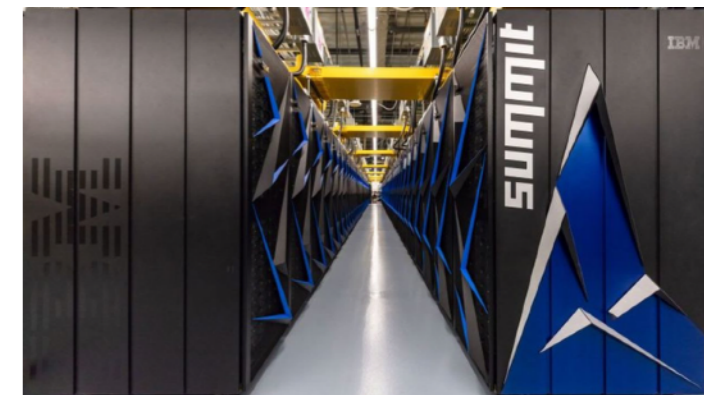
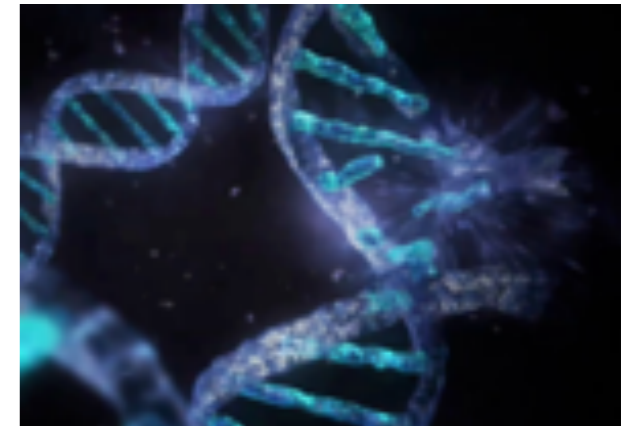
Damage Estimation for Proton Irradiation



- Primary particles
 - Protons with $E_{\text{kin}} = 500 \text{ keV} - 100 \text{ MeV}$
- Reasonable agreement with the experimental data and the PARTRAC results

Future Plans

- **Extending functionalities in MPEXS-DNA for nanodosimetry simulations**
 - More complex geometry exported from CAD
 - ex. DNA double helix structure
 - Various materials other than water (ex. DNA components)
 - Multi-GPU simulations on Cloud services / Supercomputer
 - **“Almost infinite” scalability**
 - ex. 27,648 GPUs (TESLA V100) @ Summit
 - **MPEXS-DNA will pioneer a scientific frontier in radiobiology**
 - Better understanding mechanism of DNA damages
 - More precise estimation of effects on chronic radiation exposure
 - ex. Astronauts, Airline crew, medical diagnostic, ...
 - Quicker development of next-generation radiotherapy technique
 - ex. Flash radiotherapy
 - ...
- **Further developments are continuing**



Summary

- MPEXS-DNA is a nanodosimetry simulator based on Geant4-DNA running on GPU
 - Most of functionalities in Geant4-DNA are available
 - A paper on MPEXS-DNA was published from Medical Physics
 - DOI: 10.1002/mp.13370
- Recent activities:
 - Implemented Plante's approach with a step-by-step method in Geant4-DNA and MPEXS-DNA as an option of chemistry simulation
 - Quantitative estimation of radiation damages of DNA molecules with a simpler approach
 - Simple cell model / clustering model
 - Reasonable agreement with the measurement data
 - These developments will be delivered soon in both MPEXS-DNA and Geant4-DNA
 - MPEXS-DNA also boosts the development of Geant4-DNA
- MPEXS-DNA pioneers a new promising frontier in radiobiology!

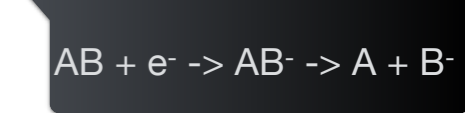
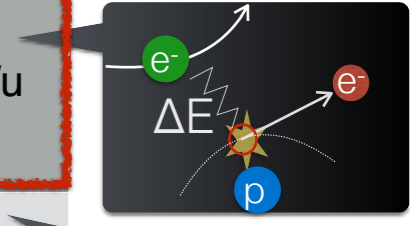
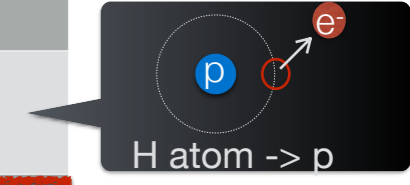
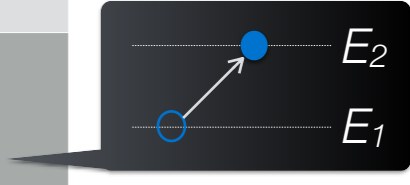
Backup

MPEXS License Model

- Two models:
 - Collaboration license
 - Only for groups who expect to submit a paper jointly within 1 year
 - Free of charge
 - The newly earned intellectual properties in the collaboration are requested to be donated to KEK or to be put in the public domain
 - Also shall not exercise the moral rights of the author or any other similar rights
 - General license
 - Contracts with KEK are required
 - A license fee will be requested

MPEXS-DNA Physics Processes

Particles		Electrons	Protons	Hydrogen atoms	Helium atoms (He ⁺⁺ , He ⁺ , He ⁰)	Li, Be, B, C, N, O, Si, and Fe
Physics Processes	Elastic scattering	9 eV - 10 keV Uehara 10 keV - 1 MeV Champion	100 eV - 1 MeV Hoang		100 eV - 10 MeV Hoang	—
	Excitation	10 eV - 10 keV Emfietzoglou 10 keV - 1 MeV Born	10 eV - 500 keV Miller Green 500 keV - 100 MeV Born	10 eV - 500 keV Miller Green	1 keV - 400 MeV Miller Green	—
	Charge change	—	100 eV - 10 MeV Dingfelder	100 eV - 10 MeV Dingfelder	1 keV - 400 MeV Dingfelder	—
	Ionization	10 eV - 10 keV Emfietzoglou 10 keV - 1 MeV Born	100 eV - 500 keV Rudd 500 keV - 100 MeV Born	100 eV - 100 MeV Rudd	1 keV - 400 MeV Rudd	0.5 MeV/u - 1.0x10 ⁶ MeV/u Rudd
	Vibrational excitation	2 - 100 eV Michaud et al.	Atomic deexcitation occurs during ionization process, and emits auger electrons and X-rays			
	Disociative attachment	4 - 13 eV Melton				



Physics Processes for X-rays (The Livermore Model)	
Compton scattering	100 eV - 1 GeV
Photoelectric effect	100 eV - 1 GeV
Gamma conversion	100 eV - 1 GeV
Rayleigh scattering	100 eV - 1 GeV