

Modeling carbon sputtering and near-surface plasma chemistry*

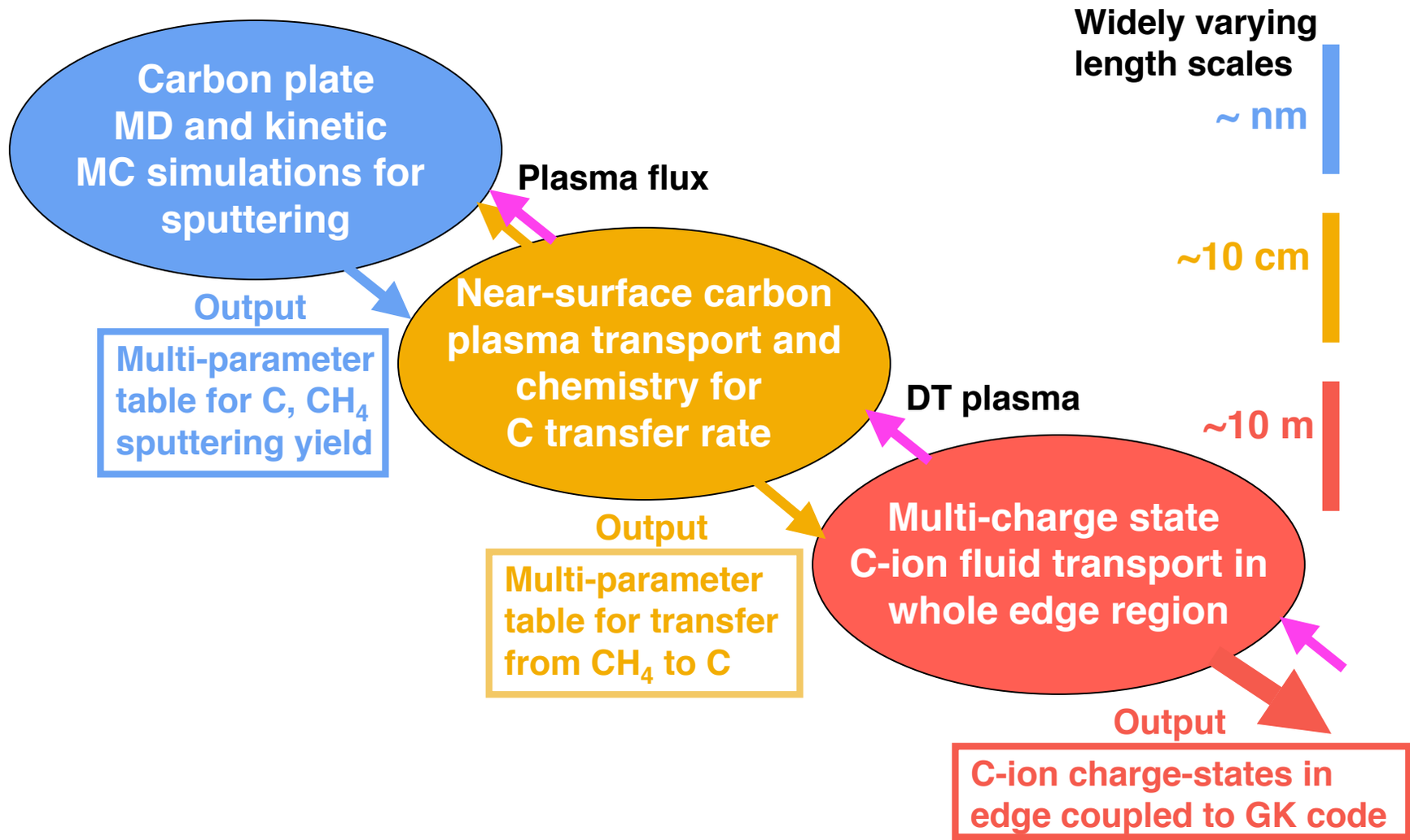
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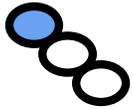
Plasma Facing Components Workshop
Princeton Plasma Physics Lab
May 9, 2005

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Modeling impurity content in fusion plasmas requires coupling 3 regions

Impurities in the edge plasma are important for power balance



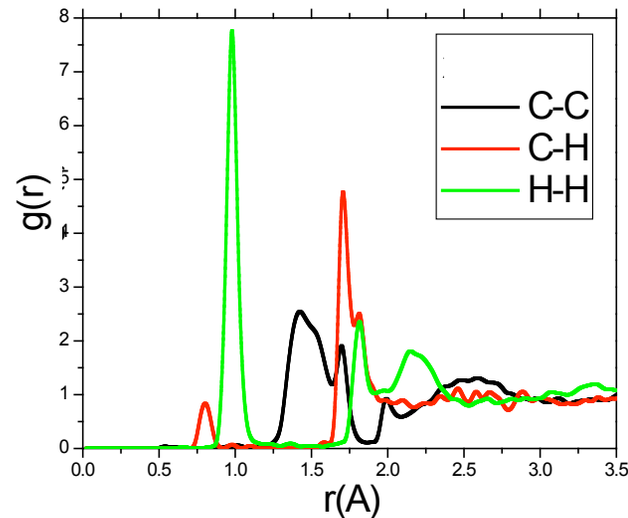
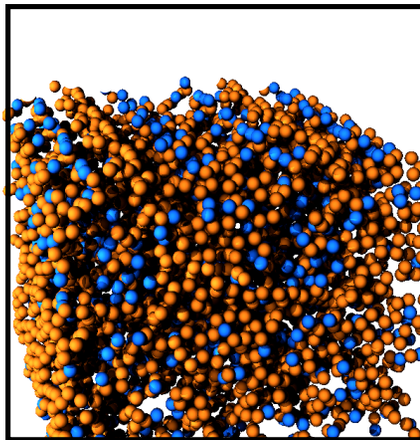


A crucial step for realistic MD simulations is proper construction of the target material

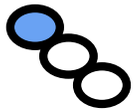
We have succeeded in developing targets corresponding to long-term exposure to reactor plasma as follows:

- start with amorphous graphite formed by pressurized melting and quenching.
- include 25% H (deuterium/tritium) in target to account for plasma exposure.
- anneal C/H target to stabilize structure.
- bombard with tritium/deuterium to include the effect of steady-state exposure.

Resulting carbon MD target showing H (blue)



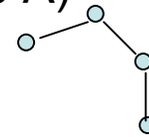
Normalized spatial distribution of lattice ions



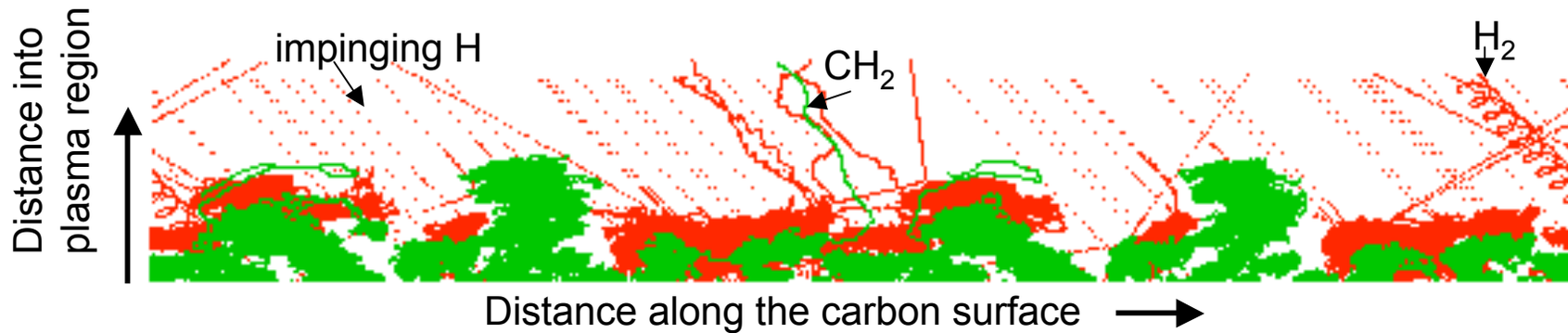
Sputtering MD simulations utilize state-of-the-art AIREBO & Brenner inter-atomic potentials

AIREBO (Adaptive Intermolecular Reactive Empirical Bond Order) potential is an extension of REBO that includes:

- Short-range, bonding interactions from Brenner ($<3 \text{ \AA}$)
- Long-range, non-bonding interactions ($<6 \text{ \AA}$)
- Torsional interactions (4-body) \longrightarrow

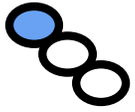


Sputtered-particle trajectories (solid colors) just above the surface



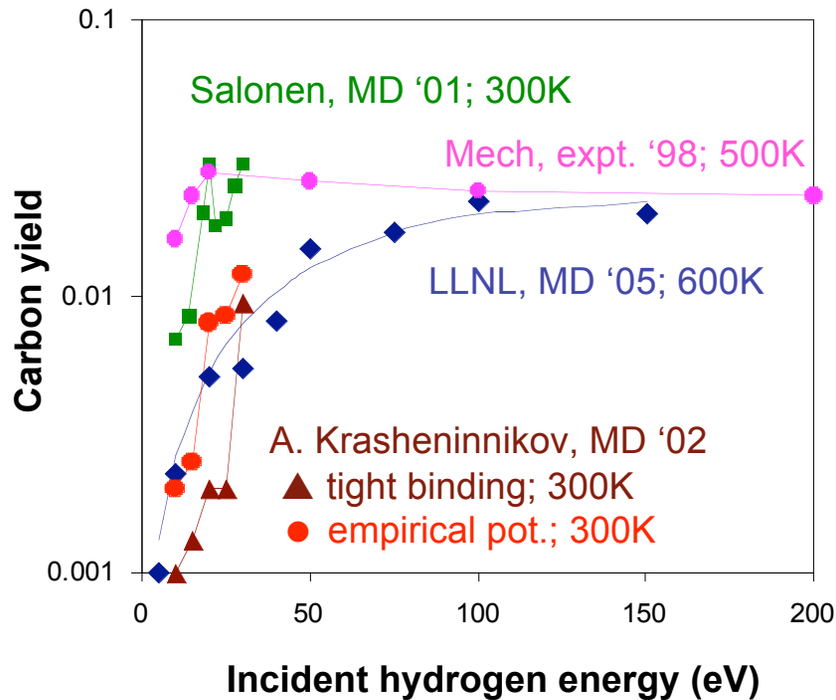
Conditions: $T_{\text{surf}} = 500\text{K}$, $E_{\text{inc}} = 20\text{eV}$, $\Theta_{\text{inc}} = 30^\circ$

After 40 D/T impacts: 6H, 2H₂, CH₂

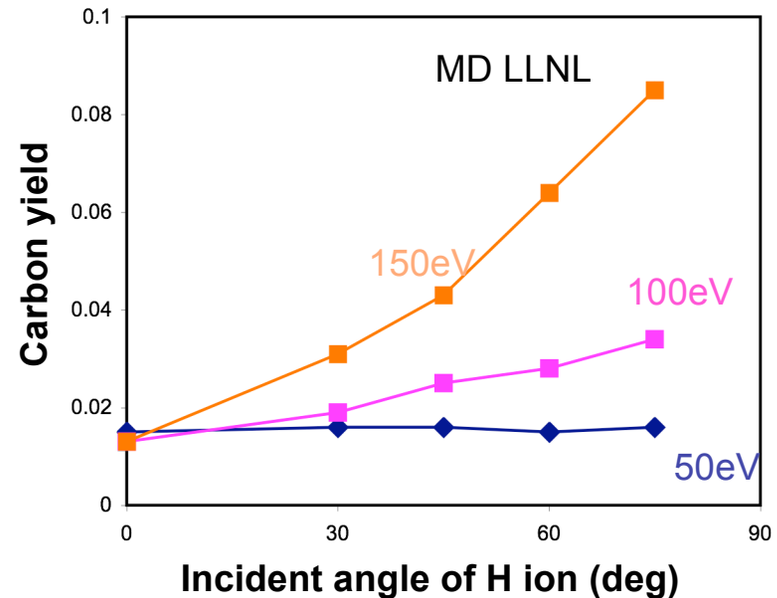


Our MD simulation results cover a range of impact energies and angles; understanding is developing

Range of results depends partly on different materials/models



Variations with angle and energy suggest impact of chemical processes

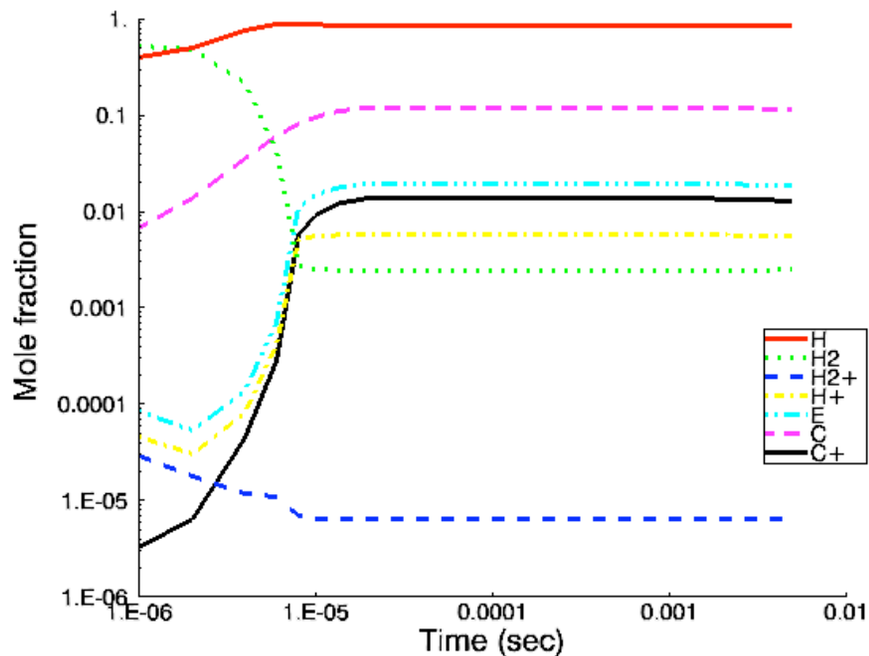


Mech, Haasz et al., J. Nucl. Mat. **255** (1998) 153
Salonen, et al., Phys. Rev. **B63** (2001) 195415
A. Krasheninnikov et al., Comp. Mat. Sci. **25** (2002) 427
LLNL, PFC and LDRD '05

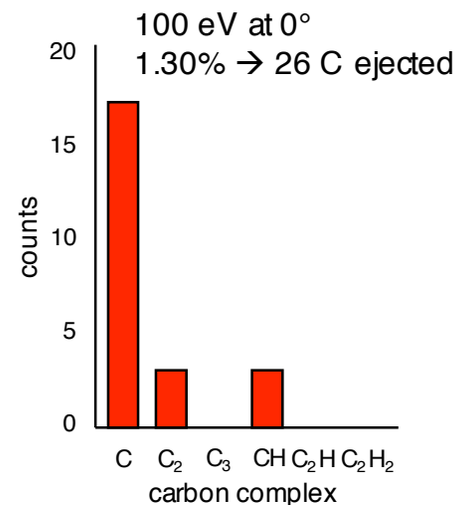
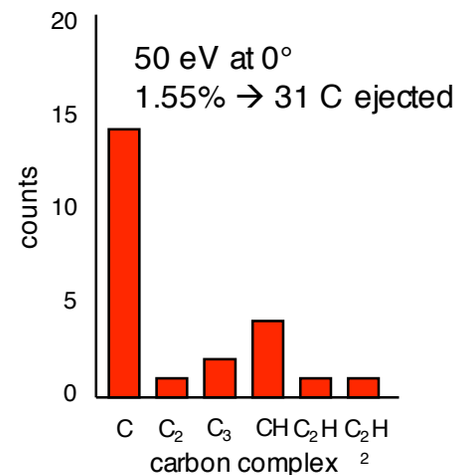


Initial reaction-diffusion model being used to rank importance of species

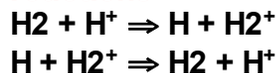
Sputtered hydrocarbons predicted by MD are now being added to find net C yield



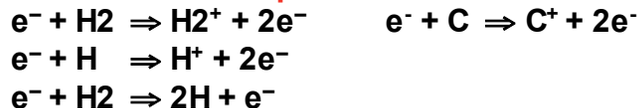
Next step:
Include sputtered hydrocarbon.



Ion-neutral reactions

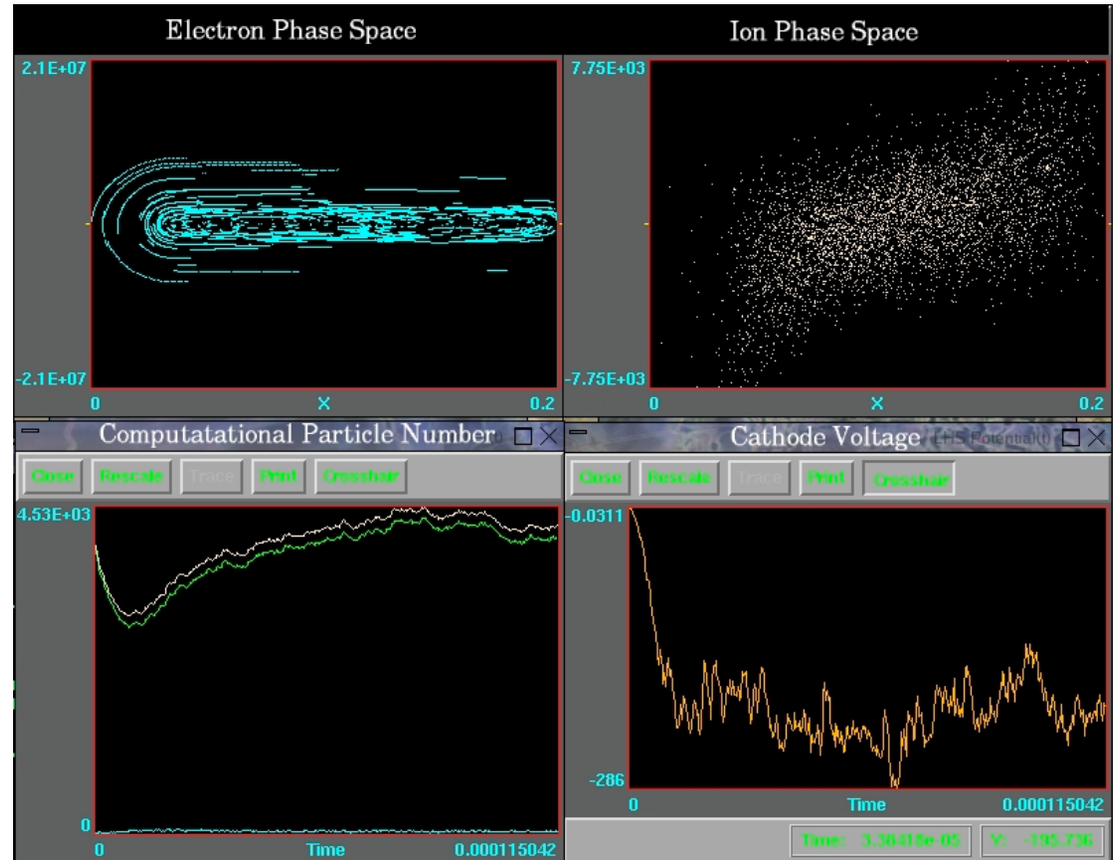


Electron impact ionization



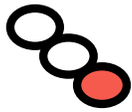
XOOPIC can use full particle electrons, but we need more efficient Boltzmann electron model

- PIC ions, Boltzmann-PIC hybrid electrons
- Electrons above specified threshold treated as particles – retains kinetic effects, Monte Carlo collision model
- Electron bulk modeled as inertialess Maxwell-Boltzmann distribution:
$$n(\mathbf{x}) = n_0 \exp(-q\phi(\mathbf{x})/T)$$
- Can choose arbitrary Boltzmann electron distribution function, $f(E)$, e.g. with cutoff tails.
- Boltzmann species collisions based on $f(E)$

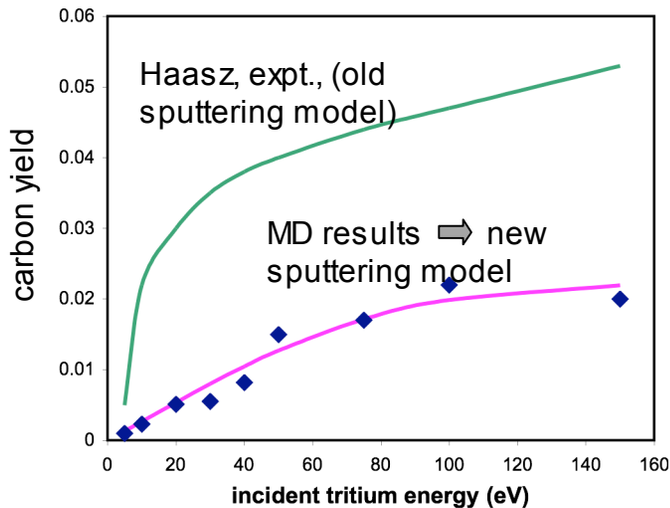
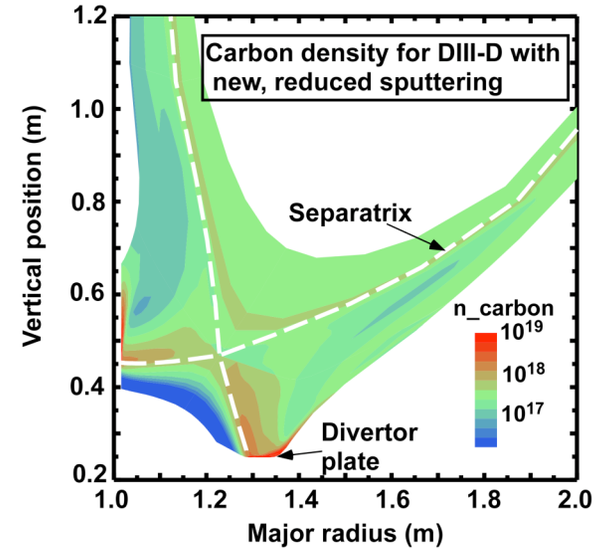
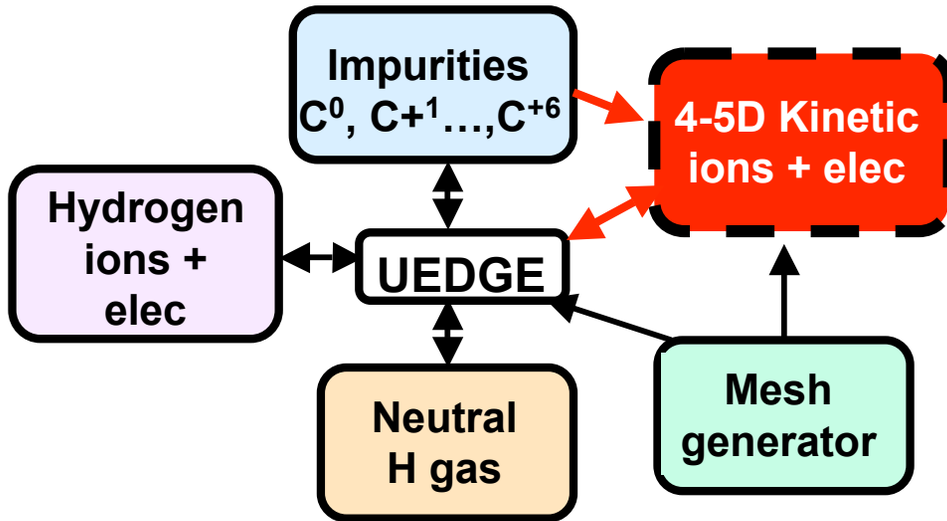


Current-driven 1D DC discharge runs up to 100 times faster than full PIC electron model.

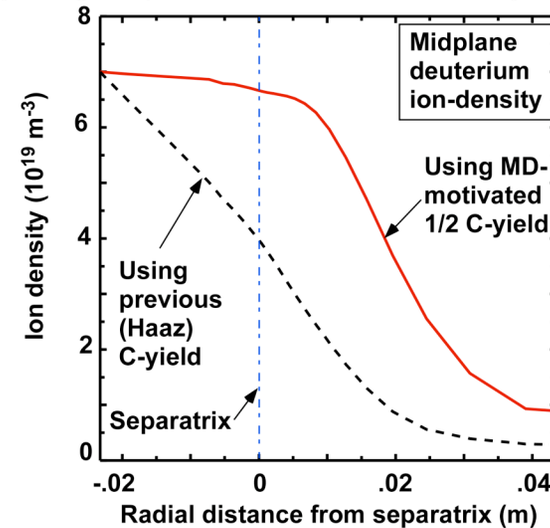
Based on Cartwright et al., *Phys. Plasmas* 7, 3252 (2000).



Full edge profile of carbon ions (charges Z= 1-6) from fluid UEDGE provides impurities

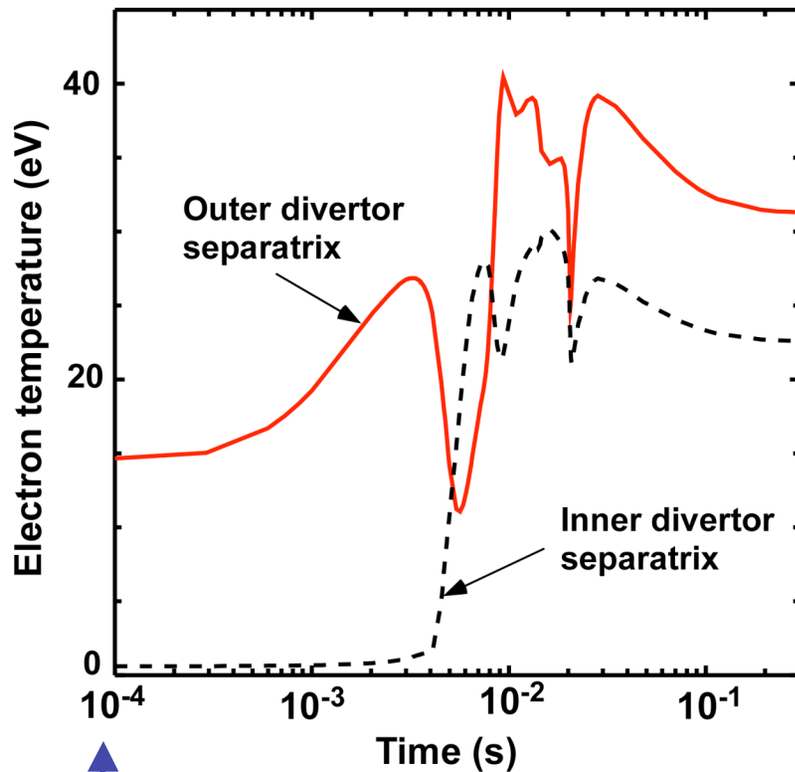


Hydrogen plasma sensitive to C-yield

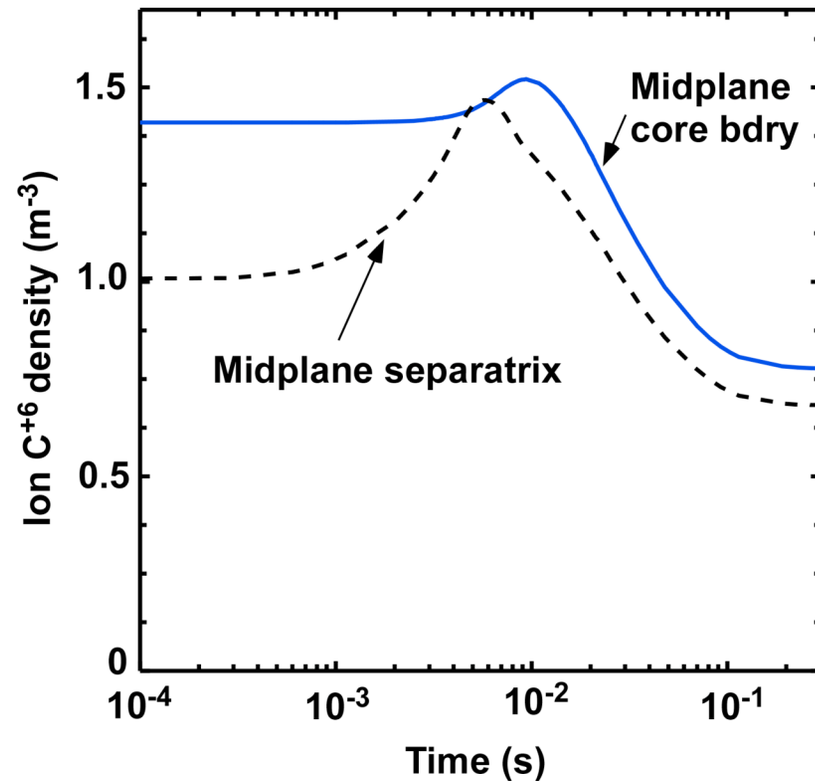


Reduced carbon sputtering induces change from “detached” inner divertor to “attached”

Divertor T_e history



Midplane C⁺⁶ history



Change from 100% to 50% of Haaz '97 sputtering model

Summary

- **Developed realistic chemically-evolved (surface roughness) C/H targets using annealing and deposition**
- **State-of-the-art carbon interaction models were applied to producing new multi-variable chemical/physical sputtering**
- **Implemented and began testing chemical-rate methodology (ChemKin) to identify dominant hydrocarbon species**
- **Initial demonstration of edge plasma sensitivity to carbon content**
- **Implemented fast Boltzmann electron model for developed near-surface dynamics via XOOPIIC plasma/neutral code**