
Particle-in-cell simulations

Part II: Implementation of Zeltron

Benoît Cerutti

CNRS & Université Grenoble Alpes, Grenoble, France.

Plan of the lectures

- **Wednesday:**

- *Morning*: The PIC method, numerical schemes and main algorithms.
- *Afternoon*: Coding practice of the Boris push and the Yee algorithm.

- **Thursday:**

- *Morning*: Implementation of Zeltron, structure and methods.
- *Afternoon*: Zeltron hands on relativistic reconnection simulations
- *Evening*: Seminar applications of PIC to relativistic magnetospheres.

- **Friday:**

- *Morning*: Boundary conditions and parallelization in Zeltron.
- *Afternoon*: Zeltron Hands on relativistic collisionless shocks simulations

General presentation

Zeltron is an **explicit, relativistic 3D PIC code** created from scratch in 2012. Originally designed to study particle acceleration in relativistic magnetic reconnection sites applied to astrophysics.

Main developers:

Benoît Cerutti (*CNRS / Univ. Grenoble Alpes*)

Greg Werner (*University of Colorado*)

Some general features



- Written in **Fortran 90**
- **Yee FDTD** algorithm for the fields
- **Boris push** for the particles
- **Efficiently parallelized** with MPI (3D domain decomposition)
- Includes **synchrotron** and **inverse Compton** radiation reaction forces
- **Non Cartesian-mesh**: spherical, cylindrical, Schwarzschild (not public)
- Large set of tools for **data reduction** and **data analysis** on the fly
- Set of **boundary conditions** (absorption, creation, open, reflective, ...)
- **No need for external libraries**

Global structure

PROGRAM:

main.f90: Contains the main body of the code, initialization, main loop in time.

MODULES :

- *mod_input.f90*: Input file where all the numerical and physical parameters can be set. This is the only file to modify for a given setup.
- *mod_initial.f90*: Initialize the fields and the particles (problem dependent).
- *mod_fields.f90*: Solve Maxwell equations and other operations related to fields.
- *mod_motion.f90*: Contains the particle push, applies boundary conditions and exchange particles at the boundaries between processors
- *mod_rhoj.f90*: For depositing particle charge and current on the grid.
- *mod_analysis.f90*: Data reduction and analysis on the fly and output to the disk
- *Others*: More technical modules related to I/O, interpolation, and parallelization.

Computation procedure per timestep in PIC

Step 1

Solve Newton's
equation

$$\frac{d\mathbf{p}}{dt} = q \left(\mathbf{E} + \frac{\mathbf{v} \times \mathbf{B}}{c} \right)$$

$$\frac{\partial \mathbf{B}}{\partial t} = -c \nabla \times \mathbf{E}$$

$$\frac{\partial \mathbf{E}}{\partial t} = c \nabla \times \mathbf{B} - 4\pi \mathbf{J}$$

Δt

Solve Maxwell's
equations (E,B)

Step 3

Deposit
Charge and
current densities
(ρ, \mathbf{J})

Step 2

General structure of the main.f90 file

INITIALISATION

1. Initialize the MPI environment for parallel computing (see *lecture III*)
2. Initialize the spatial grid, and Yee mesh (*this lecture*)
3. Set the initial conditions (Particles and fields) at $t=0$ (*this lecture*)
4. Write initial data to disk
5. Leap-frog initialization: evolve $\mathbf{u}^0 \rightarrow \mathbf{u}^{0-1/2}$ (see *lecture I*)

MAIN LOOP IN TIME

- For each particle species

1. Boris push: $\mathbf{u}^{n-1/2} \rightarrow \mathbf{u}^{n+1/2}$ (see *lecture I*)
2. Deposit currents: $\mathbf{J}_0 = 0.5\rho^n \mathbf{v}^{n+1/2}$ (see *lecture I*)
3. Evolve particle positions: $\mathbf{r}^n \rightarrow \mathbf{r}^{n+1}$
4. Apply boundary conditions and MPI communications (see *lecture III*)
5. Deposit currents: $\mathbf{J} = 0.5\rho^{n+1} \mathbf{v}^{n+1/2} + \mathbf{J}_0$ (see *lecture I*)

Sample from the main.f90 file (particles)

```
! Update u from t-dt/2 to t+dt/2, particle positions unchanged
CALL BORIS_PUSH(-1d0,me,pcl_ed,pcl_data_ed,Bxg,Byg,Bzg,Exg,Eyg,Ezg, &
               Uph,xgp,ygp,Esyned,Eicsed,NED)

! Computation of rho at t and half of the current density
CALL RHOJ(-1d0,pcl_ed,rhoed,Jx0,Jy0,Jz0,xgp,ygp,NED,id,ngh,TOPO_COMM,ierr)

! Push the particles from t to t+dt
CALL PUSH_PARTICLES(pcl_ed,NED)

! Applying boundary conditions to the particles
CALL BOUNDARIES_PARTICLES(pcl_ed,pcl_data_ed,taged,NED)

! Counting the particles leaving each subdomain
CALL COUNT_ESCAPE(pcl_ed,xminp,xmaxp,yminp,ymaxp,NED,NESC)

! Exchange of particles at the boundaries between processes
CALL COM_PARTICLES(pcl_ed,pcl_data_ed,taged,xminp,xmaxp,yminp,ymaxp, &
                  NED,NESC,id,ngh,TOPO_COMM,ierr)

! Computation of rho at t+dt and the other half of the current density
CALL RHOJ(-1d0,pcl_ed,rhoed,Jxed,Jyed,Jzed,xgp,ygp,NED,id,ngh,TOPO_COMM,ierr)

! Total current density at t+dt/2
Jxed=Jxed+Jx0
Jyed=Jyed+Jy0
Jzed=Jzed+Jz0
```

General structure of the main.f90 file

MAIN LOOP IN TIME (continues)

- For the fields

1. Collect currents from all species, and put current on the Yee mesh.
2. Push B field half time step: $B^n \rightarrow B^{n+1/2}$ (see *lecture I*)
3. Push E field full time step: $E^n \rightarrow E^{n+1}$ (see *lecture I*)
4. Correct the E field (charge conservation) with Poisson solver
(*this lecture*)
5. Push B field half time step: $B^{n+1/2} \rightarrow B^{n+1}$ (see *lecture I*)

- Analyze and write data to disk every FDUMP time steps (*this lecture*)

- Create a checkpoint every FSAVE time steps

Sample from the main.f90 file (fields)

```
!=====
! B FIELD at t=t+dt/2
!=====

CALL PUSH_BHALF (Bx, By, Bz, Ex, Ey, Ez, xgp, ygp, id, ngh, TOPO_COMM, ierr)

!=====
! SOLVE MAXWELL'S EQUATIONS at t=t+dt
!=====

! E FIELD at t=t+dt
CALL PUSH_EFIELD (Bx, By, Bz, Ex, Ey, Ez, Jx, Jy, Jz, xgp, ygp, id, ngh, TOPO_COMM, ierr)

IF (MOD(it, FREQ_POISSON) .EQ. 0) THEN
! Solve Poisson equation to ensure that  $\text{div}(\mathbf{E})=4\pi\rho$ 
CALL CORRECT_EFIELD (Ex, Ey, xgp, ygp, rho, id, ngh, TOPO_COMM, ierr)
END IF

! B FIELD at t=t+dt
CALL PUSH_BHALF (Bx, By, Bz, Ex, Ey, Ez, xgp, ygp, id, ngh, TOPO_COMM, ierr)

CALL FIELDS_NODES (Bx, By, Bz, Ex, Ey, Ez, Bxg, Byg, Bzg, Exg, Eyg, Ezg, xgp, ygp, &
                  id, ngh, TOPO_COMM, ierr)
```

The grid and the Yee mesh

- Define number of cells and spatial boundaries from the input file (*mod_input.f90*)

```
! Number of cells in X
INTEGER*8, PARAMETER, PUBLIC :: NCX=128

! Number of cells in Y
INTEGER*8, PARAMETER, PUBLIC :: NCY=128

! Spatial boundaries in the X-direction
DOUBLE PRECISION, PARAMETER, PUBLIC :: xmin=0d0, xmax=100.0

! Spatial boundaries in the Y-direction
DOUBLE PRECISION, PARAMETER, PUBLIC :: ymin=0d0, ymax=100.0

! Spatial step
DOUBLE PRECISION, PARAMETER, PUBLIC :: dx=(xmax-xmin)/NCX
DOUBLE PRECISION, PARAMETER, PUBLIC :: dy=(ymax-ymin)/NCY
```

- Build the **spatial arrays** (*main.f90*)

```
! Nodal lattice
DO ix=1,NX
  xg(ix)=(ix-1)*1d0/((NX-1)*1d0)*(xmax-xmin)+xmin
ENDDO

DO iy=1,NY
  yg(iy)=(iy-1)*1d0/((NY-1)*1d0)*(ymax-ymin)+ymin
ENDDO

! Yee lattice
xyee=xg+dx/2.0
yyee=yg+dy/2.0
```


Particle data structure

The particle distribution function is characterized by 3 arrays (*mod_input.f90*):

```
! BACKGROUND ELECTRONS distribution function components
DOUBLE PRECISION, ALLOCATABLE, PUBLIC :: pcl_eb(:, :)
DOUBLE PRECISION, ALLOCATABLE, PUBLIC :: pcl_data_eb(:, :)
INTEGER*8, ALLOCATABLE, PUBLIC      :: tageb(:)
```

ALLOCATE (pcl_eb(1:7, 1:NP)) : Actual particle distribution function (pb independent)


List of particles



	x	y	z	u _x	u _y	u _z	wgt
Part 1							
Part 2							
...							
Part NP							

ALLOCATE (pcl_data_eb(1:4, 1:NP)) : Additional particle data information (pb dependent)

List of particles



	Data 1	Data 2	Data 3	Data 4
Part 1				
Part 2				
...				
Part NP				


Particle data structure

The particle distribution function is characterized by 3 arrays (*mod_input.f90*):

```
! BACKGROUND ELECTRONS distribution function components
DOUBLE PRECISION, ALLOCATABLE, PUBLIC :: pcl_eb(:, :)
DOUBLE PRECISION, ALLOCATABLE, PUBLIC :: pcl_data_eb(:, :)
INTEGER*8, ALLOCATABLE, PUBLIC      :: tageb(:)
```

ALLOCATE (tag_eb (1:NP)) : Each particle is identified by a **unique integer number**
=> *Useful for particle tracking*

List of particles



	tag
Part 1	tag1
Part 2	tag2
...	...
Part NP	tag3

Fields/Currents data structure

- **Fields** defined on the Yee lattice (*main.f90*): Used to **evolve the fields** (see *Lecture I*)

```
!*****  
! Magnetic and Electric fields components Yee lattice  
DOUBLE PRECISION, DIMENSION(1:NX,1:NY) :: Bx,By,Bz  
DOUBLE PRECISION, DIMENSION(1:NX,1:NY) :: Ex,Ey,Ez
```

- The **current** density is needed on the Yee mesh (*main.f90*)

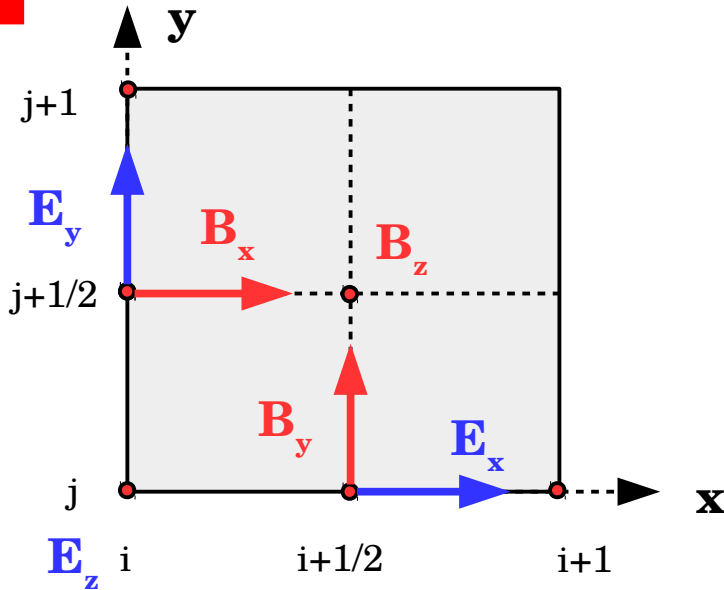
```
!*****  
! Current density components Yee lattice  
DOUBLE PRECISION, DIMENSION(1:NX,1:NY) :: Jx,Jy,Jz
```

- **Fields** defined on the nodes of the grid (*main.f90*): Used to **evolve the particles** (see *Lecture I*)

```
!*****  
! Magnetic and Electric fields components at nodes  
DOUBLE PRECISION, DIMENSION(1:NX,1:NY) :: Bxg,Byg,Bzg  
DOUBLE PRECISION, DIMENSION(1:NX,1:NY) :: Exg,Eyg,Ezg
```

From Yee to nodes

2D



- Electric field

$$Exg_{i,j} = \frac{Ex_{i+1/2,j} + Ex_{i-1/2,j}}{2}$$

$$Eyg_{i,j} = \frac{Ey_{i,j+1/2} + Ey_{i,j-1/2}}{2}$$

$$Ezg_{i,j} = Ez_{i,j}$$

- Magnetic field

$$Bxg_{i,j} = \frac{Bx_{i,j+1/2} + Bx_{i,j-1/2}}{2}$$

$$Byg_{i,j} = \frac{By_{i+1/2,j} + By_{i-1/2,j}}{2}$$

$$Bzg_{i,j} = \frac{Bz_{i+1/2,j+1/2} + Bz_{i-1/2,j+1/2} + Bz_{i+1/2,j-1/2} + Bz_{i-1/2,j-1/2}}{4}$$

Correction of the electric field

Zeltron does not use a charge conserving deposition scheme

=> **Poisson equation must be solved!**

Consider that: $\vec{E}' = \vec{E} + \delta \vec{E}$

Correct Known field Small correction

$$\delta \vec{E} = -\nabla \delta \phi \quad \longrightarrow \quad \nabla^2 \delta \phi = -(4\pi\rho - \nabla \cdot \vec{E})$$

Solve this equation using a Gauss-Seidel iterative method

“Empirically”, Zeltron uses **500 iterations**, every **25 time steps**.

Relevant parameters in *mod_input.f90*

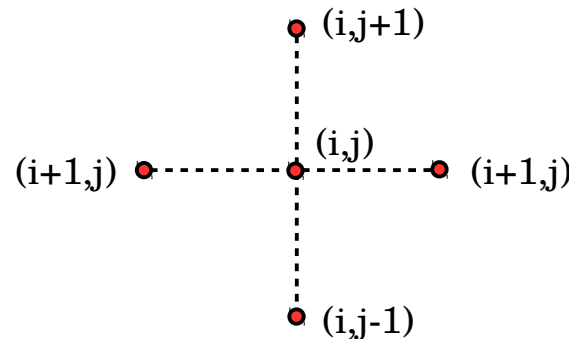
```
! Poisson solver calling frequency in terms of timesteps
INTEGER, PARAMETER, PUBLIC :: FREQ_POISSON=25

! Number of iterations to solve Poisson's equation
INTEGER, PARAMETER, PUBLIC :: NIT=500
```

Poisson solver

$$\nabla^2 \delta \phi = -(4 \pi \rho - \nabla \cdot \vec{E})$$

2D Example: 5 points stencil



$$\nabla^2 \delta \phi = \frac{\partial^2 \delta \phi}{\partial x^2} + \frac{\partial^2 \delta \phi}{\partial y^2} \approx \frac{\delta \phi_{i+1,j} - 2 \delta \phi_{i,j} + \delta \phi_{i-1,j}}{\Delta x^2} + \frac{\delta \phi_{i,j+1} - 2 \delta \phi_{i,j} + \delta \phi_{i,j-1}}{\Delta y^2}$$

Injecting this into Poisson and after some rearrangements yields (and if

$\Delta x = \Delta y$):

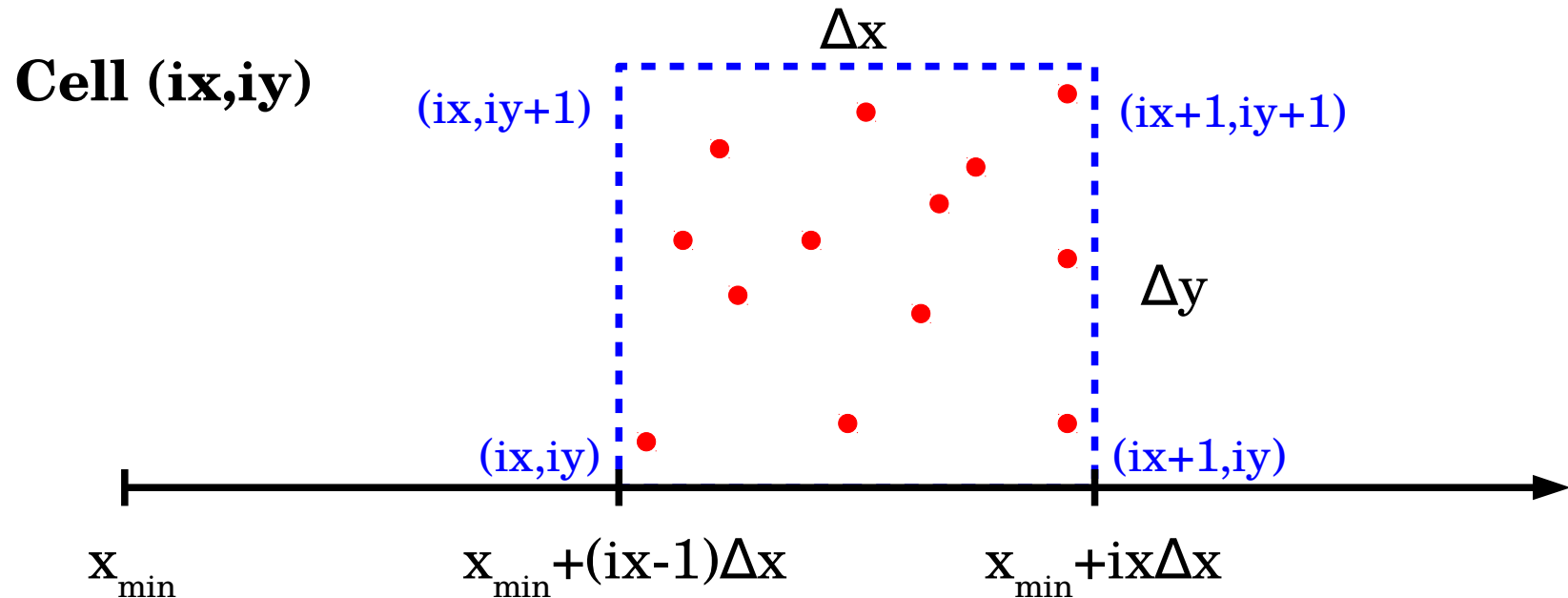
$$\delta \phi_{i,j} = \frac{1}{4} \left(\delta \phi_{i+1,j} + \delta \phi_{i-1,j} + \delta \phi_{i,j+1} + \delta \phi_{i,j-1} + (4 \pi \rho - \nabla \cdot \vec{E}) \Delta x^2 \right)$$

Calling the subroutine in *main.f90*

```
IF (MOD(it,FREQ_POISSON).EQ.0) THEN
! Solve Poisson equation to ensure that div(E)=4*pi*rho
CALL CORRECT_EFIELD (Ex, Ey, xgp, ygp, rho, id, ngh, TOPO_COMM, ierr)
END IF
```


Particle initialization: Spatial distribution

Let's consider a **uniform distribution** in 2D (*mod_initial.f90*)



```
!*****  
! Definition of the initial position (x0,y0) as a uniform  
random number defined between 0 and 1
```

```
x0c=0.0  
y0c=0.0
```

```
CALL RANDOM_NUMBER(x0c)  
CALL RANDOM_NUMBER(y0c)
```

```
x0c=xmin+(ix-1)*dx+x0c*dx  
y0c=ymin+(iy-1)*dy+y0c*dy
```

```
!*****
```

Particle weight

- A macroparticle represents a **large number** of physical particles following the exact same trajectory in phase space.

=> The particle weight gives the **normalization factor** to connect between numerical and physical plasma densities.

- For a **uniform physical** and **numerical** plasma density, all the particles have the same weight given by:

$$\text{weight} = \frac{N_{phys}}{N_{num}} = \underbrace{n}_{\text{Physical density}} \times \frac{(L_x L_y L_z)}{N_{num}}$$

Sample code in 2D from *main.f90*:

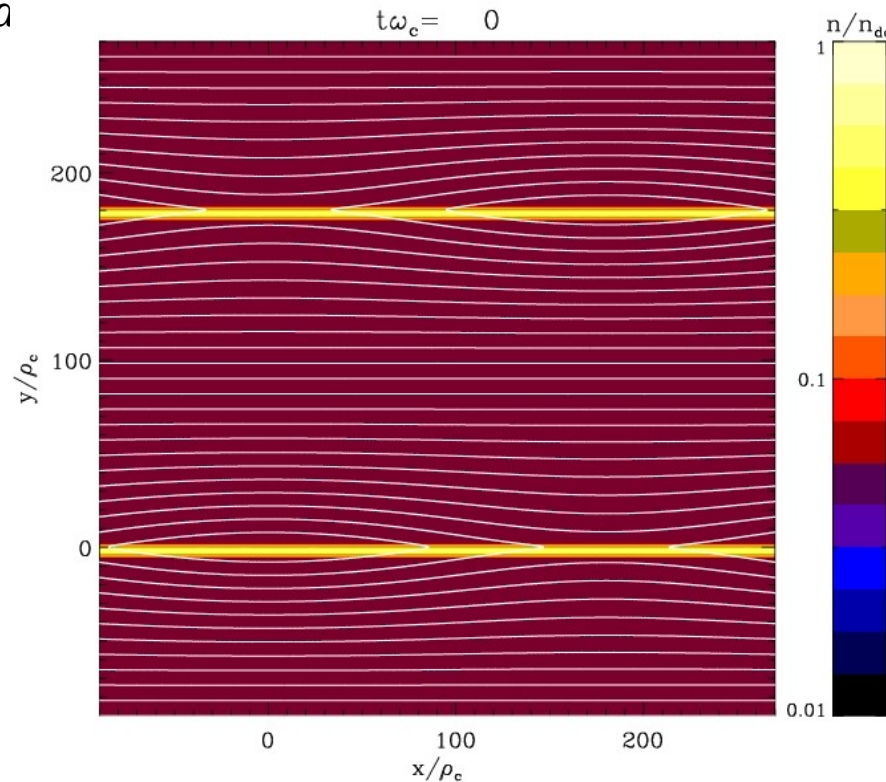
```
! Weight of background particles
pcl_eb(7, :) = density_ratio * nd0 * (xmax - xmin) * (ymax - ymin) / NP
pcl_pb(7, :) = density_ratio * nd0 * (xmax - xmin) * (ymax - ymin) / NP
```

Variable particle weight

Large **density contrast** or **sharp density profiles** are hard to model with a constant particle weight.

=> Result in a **bad sampling** of low density regions

Example: *Relativistic reconnection studies where factor >10 between the background and the sheet plasma*



Solution: Variable particle weighting: $weight(y_i) = n(y_i) \times \frac{L_x L_y L_z}{N_{num}}$

Macroscopic quantities reconstruction

One can reconstruct global and fluid quantities from the particles: (*mod_analysis.f90*)

- **Particle spectrum:** $\frac{dN}{d\gamma} \approx \frac{1}{\Delta\gamma} \sum_{k=1}^{N_{cell}} w_k$ SUBROUTINE SPECTRUM_ANGULAR

- **Plasma density:** $\frac{dN}{dV} \approx \frac{1}{\Delta V} \sum_{k=1}^{N_{cell}} w_k$ SUBROUTINE MAP_XY

- **Stress-energy tensor:** $T^{\mu\nu} = \left(\rho + \frac{P}{c^2} \right) V^\mu V^\nu - P \eta^{\mu\nu}$ SUBROUTINE MAP_FLUID

- *Energy density:* $T^{00} \equiv U_e \approx \frac{1}{\Delta V} \sum_{k=1}^{N_{cell}} w_k \gamma_k m c^2$

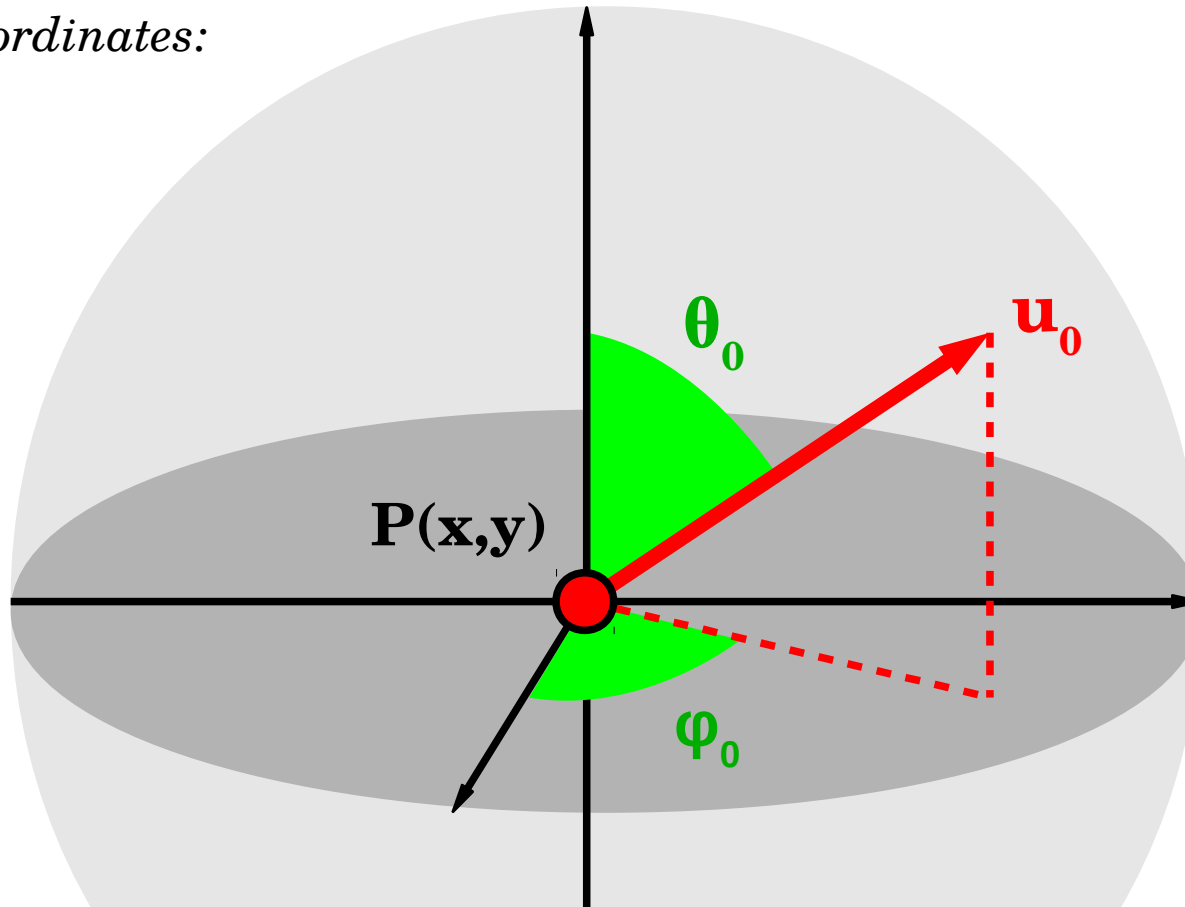
- *Momentum density:* $T^{0i} \equiv \vec{U}_p \approx \frac{1}{\Delta V} \sum_{k=1}^{N_{cell}} w_k \vec{u}_k m c$

- *Pressure tensor:* $T^{ij} \equiv P^{ij} \approx \frac{1}{\Delta V} \sum_{k=1}^{N_{cell}} \left(\frac{w_k m c^2}{\gamma_k} \right) u_k^i u_k^j$

Particle initialization: Angular distribution

Let's consider an **isotropic angular distribution** in 3D

In spherical coordinates:



- $\cos\theta_0$ uniform random number between **-1** and **1**.
- φ_0 uniform random number between **0** and **2π** .

Particle initialization: Angular distribution

Let's consider an **isotropic angular distribution** in 3D (*mod_initial.f90*)

```
!*****  
! Definition of the initial phi0 as a uniform random value  
between 0 and 1  
  
phi0=0.0  
  
CALL RANDOM_NUMBER(phi0)  
  
phi0=phi0*2.0*pi  
  
! Definition of the initial cth0=cos(theta0) as a uniform  
random value between 0 and 1  
  
cth0=0.0  
  
CALL RANDOM_NUMBER(cth0)  
  
cth0=cth0*2.0-1.0  
  
! Initial 4-velocity components  
ux0c=u0*sqrt(1.0-cth0*cth0)*cos(phi0)  
uy0c=u0*sqrt(1.0-cth0*cth0)*sin(phi0)  
uz0c=u0*cth0
```

Particle initialization: Energy distribution

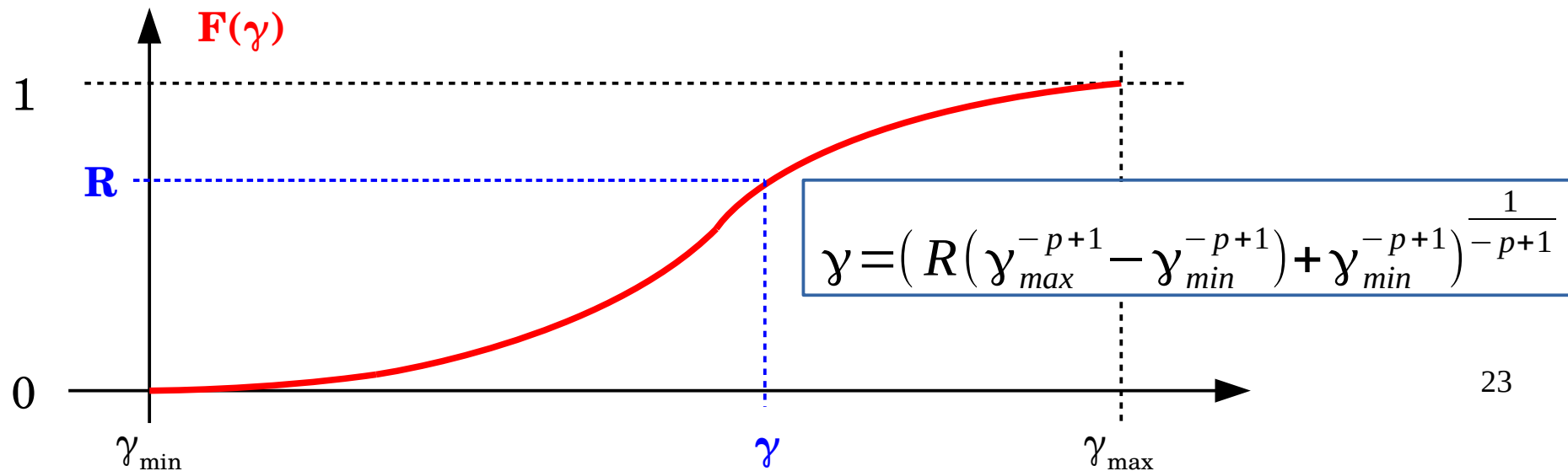
Let's consider a **Power-law distribution**:

$$f(\gamma) = \frac{dN}{d\gamma} \propto \gamma^{-p} \quad \text{with} \quad \gamma \in [\gamma_{\min}, \gamma_{\max}]$$

The trick is to use the **cumulative distribution**:

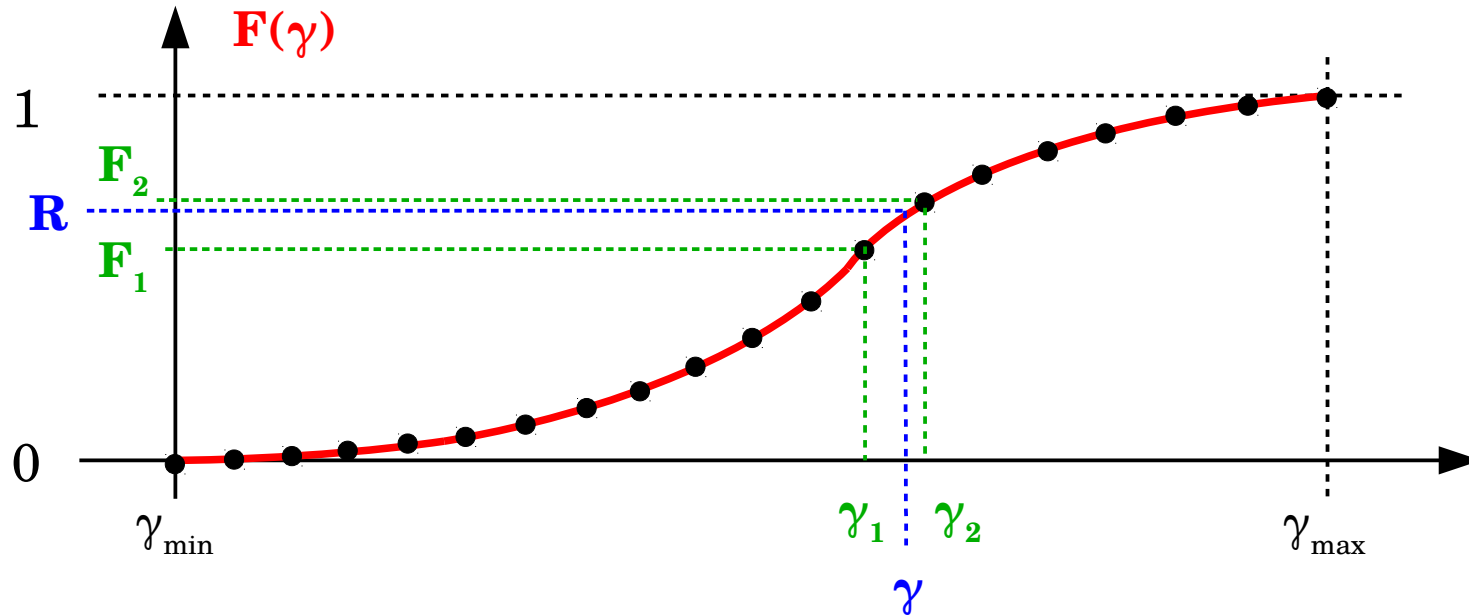
$$F(\gamma) \equiv \frac{\int_{\gamma_{\min}}^{\gamma} f(\gamma') d\gamma'}{\int_{\gamma_{\min}}^{\gamma_{\max}} f(\gamma') d\gamma'} = \frac{\gamma^{-p+1} - \gamma_{\min}^{-p+1}}{\gamma_{\max}^{-p+1} - \gamma_{\min}^{-p+1}}$$

Next, define a **uniform random number R between 0 and 1**, and sample F to invert the distribution:



Particle initialization: Energy distribution

However, the cumulative distribution is **often a tabulated** (no analytical expression)



We need to interpolate, a **linear interpolation** usually suffices:

$$R \approx (1 - \Gamma) F_1 + \Gamma F_2 \quad \Gamma = \frac{\gamma - \gamma_1}{\gamma_2 - \gamma_1}$$

$$\gamma \approx \frac{R(\gamma_2 - \gamma_1) - (\gamma_2 F_1 - \gamma_1 F_2)}{F_2 - F_1}$$

Particle initialization: Energy distribution

In Zeltron, it looks like this (*mod_initial.f90*):

```
Ru=0.0

CALL RANDOM_NUMBER (Ru)
Ru=Ru*0.9999

! Sum over all particles in a the cell
DO ic=1,PPC

! Localize closest known values of cumulative distribution gFu
minu=minloc(abs(gFu-Ru(ic)))
iu=minu(1)

! F1 and F2
gFu1=gFu(iu)
gFu2=gFu(iu+1)

! u1 and u2
u1=ud(iu)
u2=ud(iu+1)

! Linear interpolartion in the log-linear plane (numerically
more accurate)
u0(ic)=exp((Ru(ic)*(log(u2)-log(u1))-
log(u2)*gFu1+log(u1)*gFu2)/(gFu2-gFu1))

ENDDO
```

Hands-on II: Relativistic magnetic reconnection

$$\sigma = \frac{B_0^2}{4\pi n m c^2}$$

Relativistic reconnection: $\sigma \gg 1$

Relativistic Alfvén speed $V_A = c \sqrt{\frac{\sigma}{1+\sigma}} \approx c$

Dissipation of magnetic energy => relativistic particles !

Hands-on II: Relativistic magnetic reconnection

$$\sigma = \frac{B_0^2}{4\pi n m c^2}$$

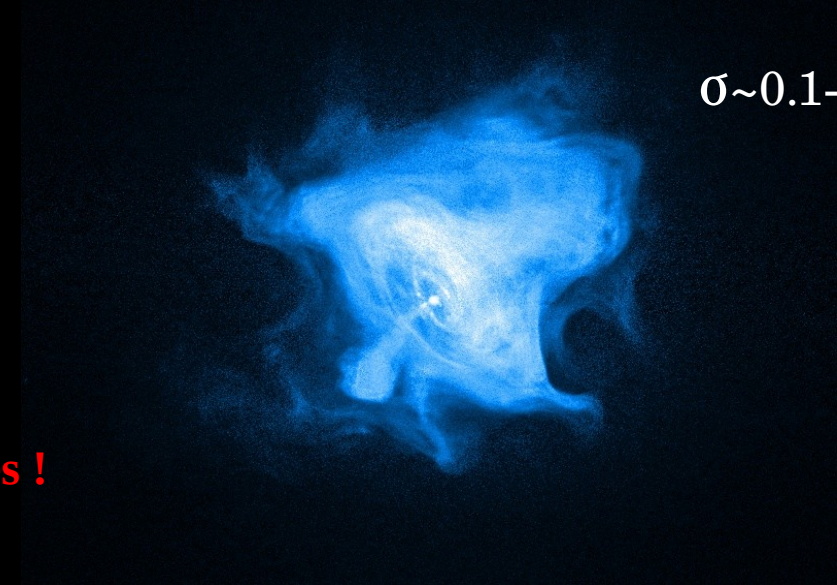
Relativistic reconnection: $\sigma > \sim 1$

Relativistic Alfvén speed $V_A = c \sqrt{\frac{\sigma}{1+\sigma}} \approx c$

Dissipation of magnetic energy => relativistic particles !

Possible astrophysical applications:

Pulsar Wind Nebulae (Crab flares)



$\sigma \sim 0.1-1$

Hands-on II: Relativistic magnetic reconnection

$$\sigma = \frac{B_0^2}{4\pi n m c^2}$$

Relativistic reconnection: $\sigma > \sim 1$

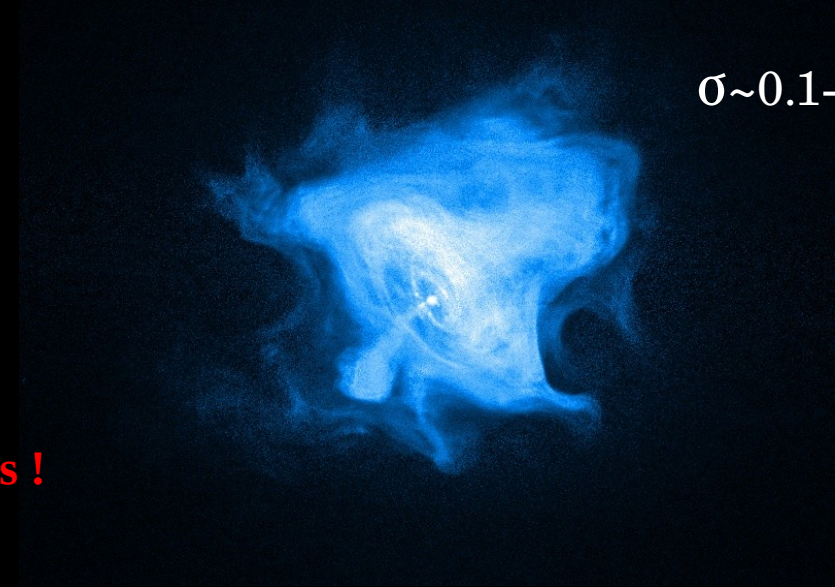
Relativistic Alfvén speed $V_A = c \sqrt{\frac{\sigma}{1+\sigma}} \approx c$

Dissipation of magnetic energy => relativistic particles !

Relativistic magnetospheres
(pulsars, magnetars, Kerr black holes)

Possible astrophysical applications:

Pulsar Wind Nebulae (Crab flares)



Hands-on II: Relativistic magnetic reconnection

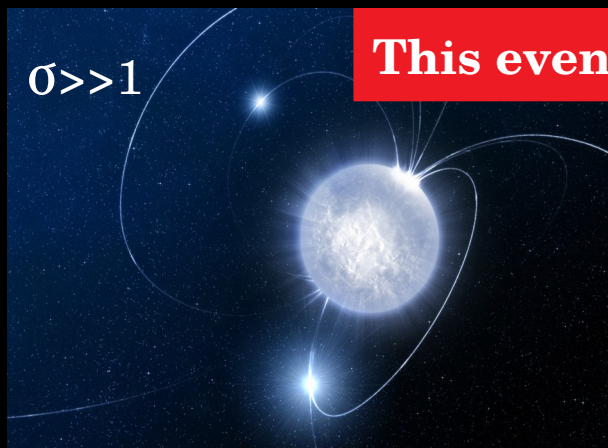
$$\sigma = \frac{B_0^2}{4\pi n m c^2}$$

Relativistic reconnection: $\sigma > \sim 1$

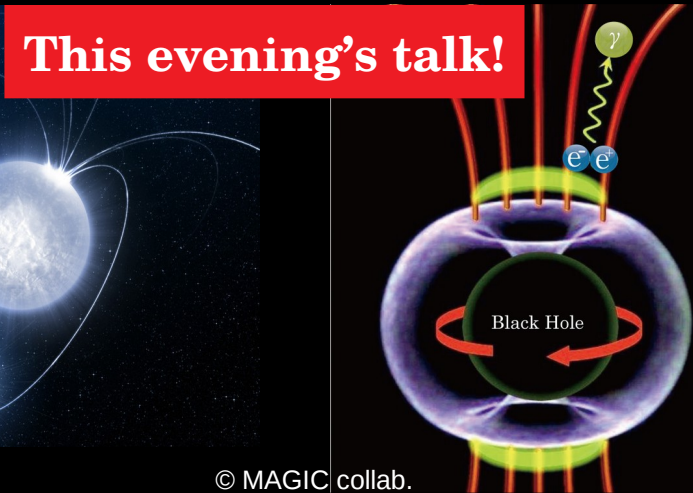
Relativistic Alfvén speed $V_A = c \sqrt{\frac{\sigma}{1+\sigma}} \approx c$

Dissipation of magnetic energy => relativistic particles !

Relativistic magnetospheres
(pulsars, magnetars, Kerr black holes)

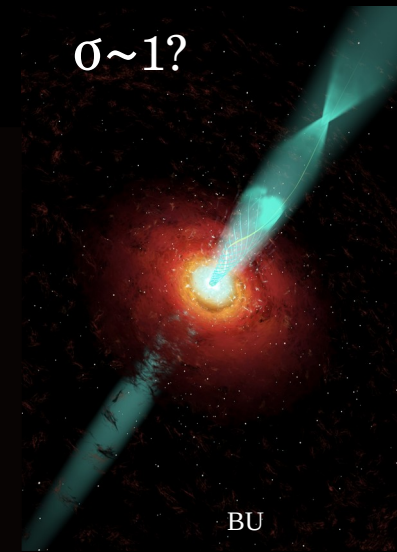
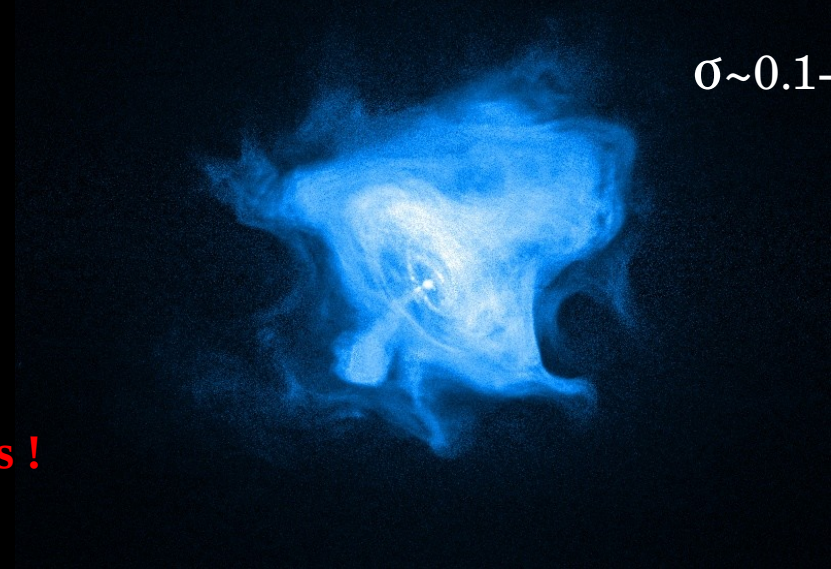


© ESO/L. Calçada

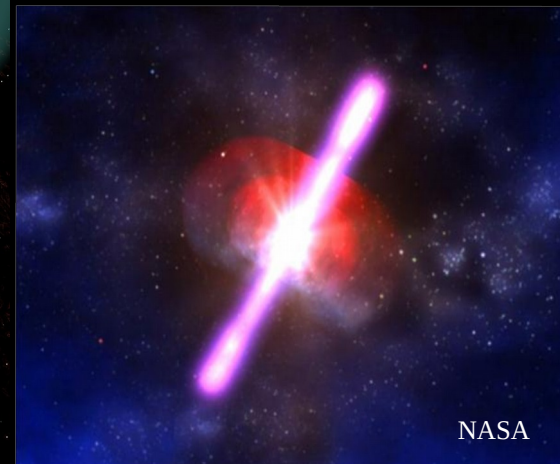


Possible astrophysical applications:

Pulsar Wind Nebulae (Crab flares)



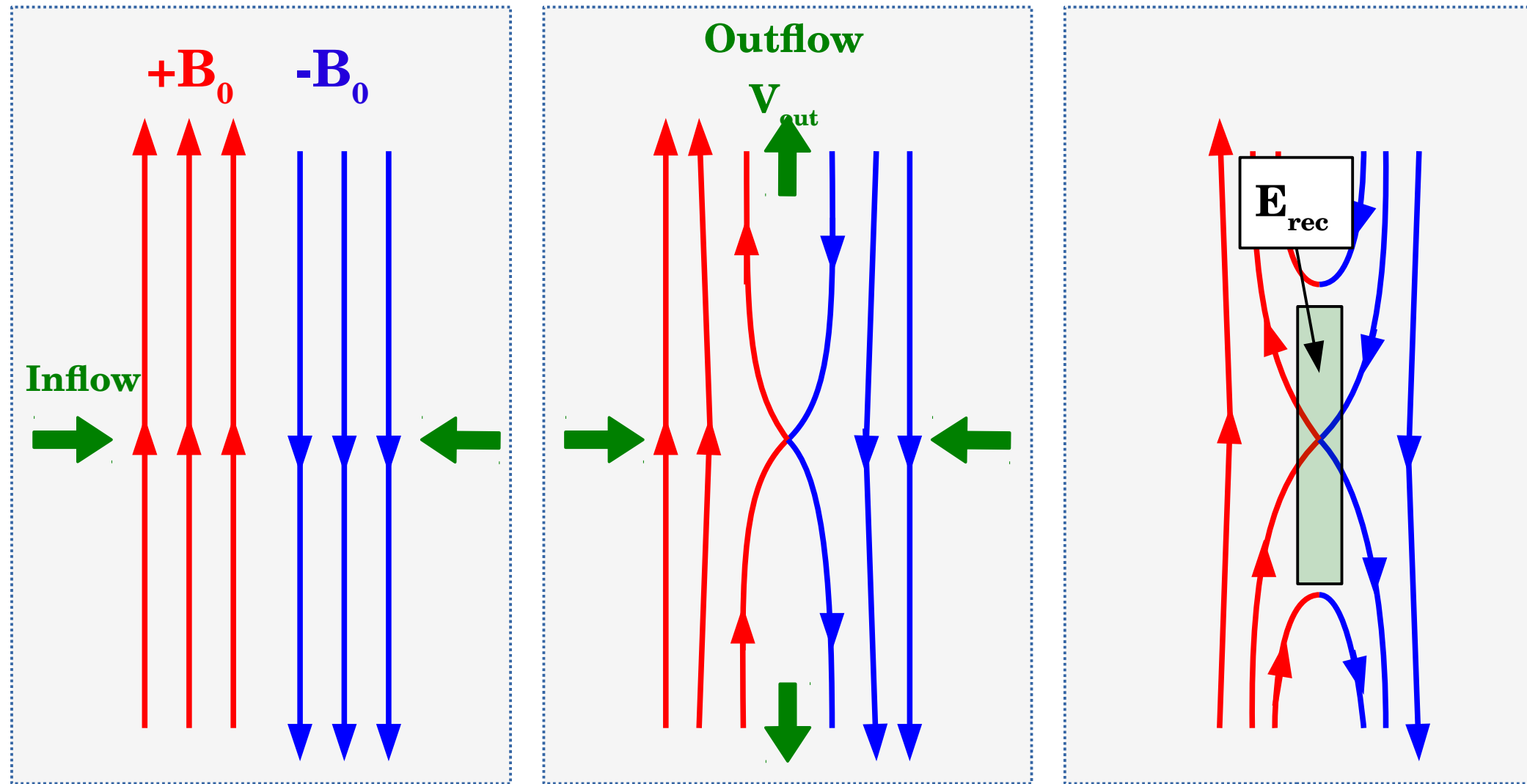
Relativistic jets



Gamma-ray burst
(prompt?)

Reconnection : A change of magnetic topology

[See reviews by Zweibel & Yamada, 2009, Kagan et al. 2015]

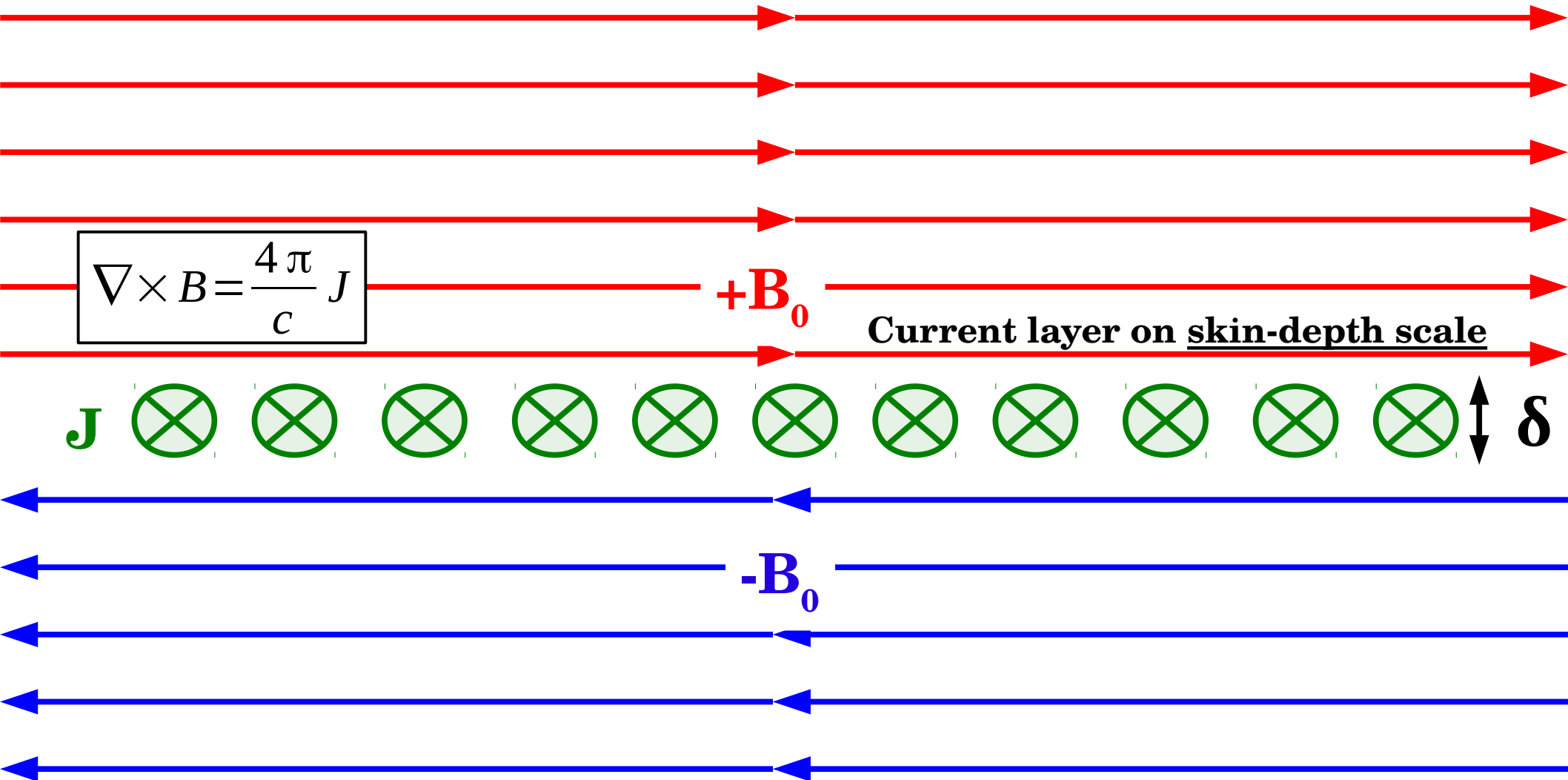


How fast does reconnection proceed?

$$\text{Reconnection rate: } \beta_{rec} = V_{in} / V_{out}$$

Magnetic energy \Rightarrow **Plasma kinetic energy** (heating+non-thermal particles)

Usual PIC setup: The Harris sheet

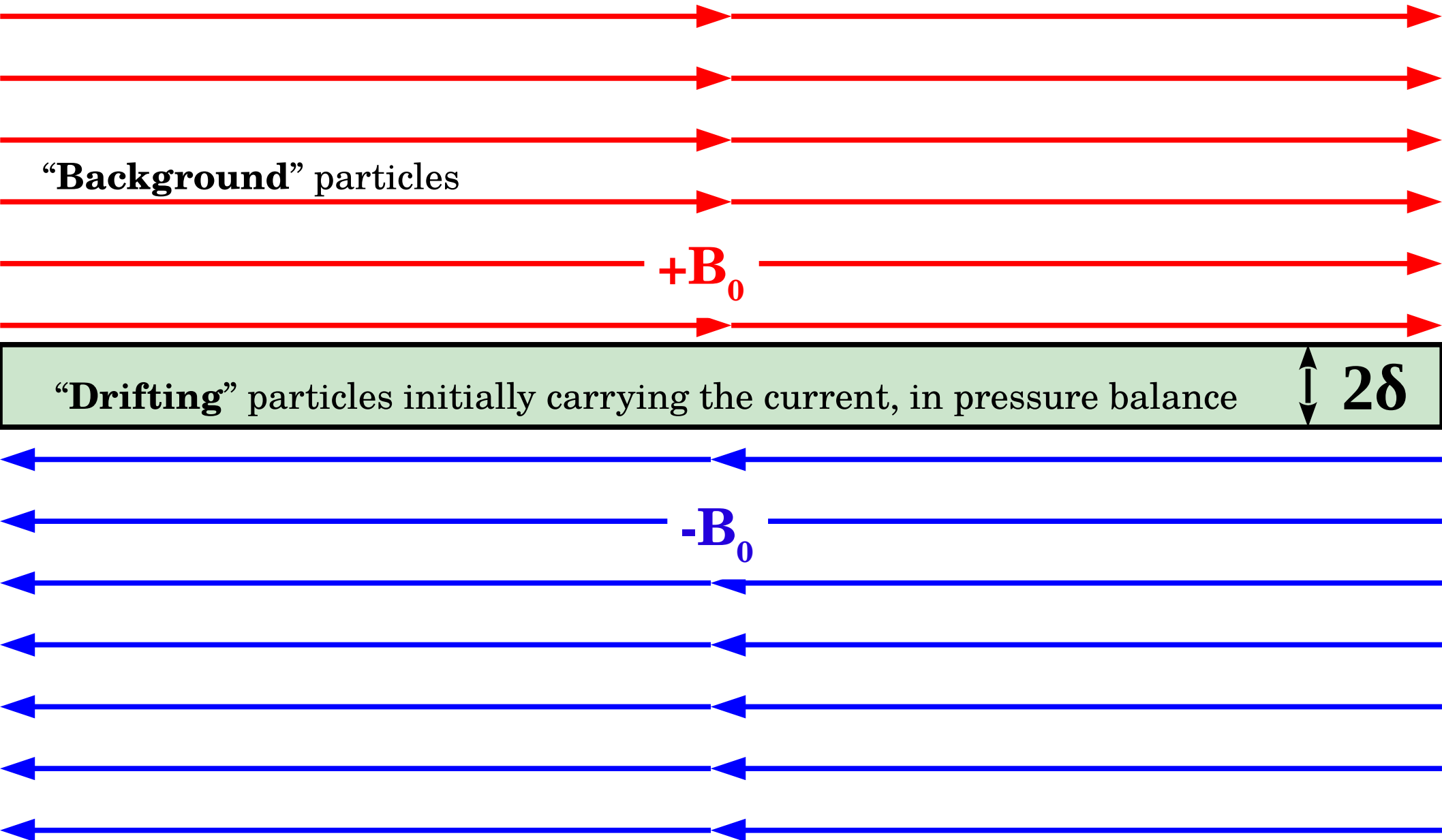


Other setup investigated :

- « **Force-Free** » current sheet (e.g., *Guo et al. 2015*)

- « **ABC** » equilibrium setup **see Journal club** (*Nalewajko+2016 ; Lyutikov+2016*)

Usual PIC setup: The Harris sheet



The relativistic Harris solution

[See Kirk & Skjæraasen 2003]

The Harris solution is a **kinetic equilibrium** between the upstream **magnetic pressure** and a **hot plasma** concentrated inside the current sheet.

The solution:

The **reconnecting** field: $\mathbf{B} = B_0 \tanh\left(\frac{y}{\delta}\right) \mathbf{e}_x$

The “**drifting**” plasma density: $n_d = n_0 \cosh^{-2}\left(\frac{y}{\delta}\right)$ $n_0 = \frac{kT}{4\pi e^2 \Gamma_d \beta_d^2 \delta^2}$

Pressure balance condition across the sheet: $\frac{B_0^2}{8\pi} = n_d kT$

The “**background**” plasma: $n_b = \epsilon n_0$, ($\epsilon \ll 1$ for $\sigma \gg 1$)

$$\sigma = \frac{B_0^2}{4\pi n_b m_e c^2} \gg 1$$

Particle initialization: Drifting Maxwellian

The **Harris** solution assumes a relativistic **drifting Maxwellian** distribution, i.e., the plasma has a relativistic motion of **Lorentz factor Γ_d** .

In the lab frame: $f(\vec{u}) \propto \exp\left(\frac{-(\gamma' - 1)}{\theta}\right)$

Where $\theta = \frac{kT'}{mc^2}$ $\gamma' = \Gamma_d(\gamma - \vec{\beta}_d \cdot \vec{u})$

Following *Swisdak (2013)*:

$$\vec{u} = \vec{u}_{\parallel} + \vec{u}_{\perp}$$



Generation of \mathbf{u}_{\parallel} via the cumulative distribution of $F(\mathbf{u}_{\parallel})$



\mathbf{u}_{\perp} is generated, but cannot be chosen independently of \mathbf{u}_{\parallel}

The full procedure is in *mod_initial.f90*

Subroutines: `SET_DRIFT_MAXWELLIAN`, `INIT_DRIFT_MAXWELLIAN`, `GEN_UP`, `GEN_PS`

Initial numerical setup

e⁺/e⁻ pairs

Time step=0, Species=ions

