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### A-Coefficient Study

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Research Notes

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# A-Coefficient Study

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### Abstract

This is a study of the asymptotic behavior of the  $\mathcal{A}$ -coefficients from Clog/acoeff.f90. Each figure is constructed by a Fortran driver grXXX.f90 and a corresponding plotting routine grXXX.sm. The plasma under consideration is equimolar DT with electron number density  $n_e$ , electron temperature  $T_e$ , and ion temperature  $T_I$ . The code used to generate the data has now been fully checked.

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### I. A-COEFFICIENTS AS A FUNCTION OF ENERGY

The A-coefficients take the form

$$\mathcal{A}_b = \mathcal{A}_b^{\text{C}} + \mathcal{A}_b^{\Delta Q} \quad \text{where} \quad \mathcal{A}_b^{\text{C}} = \mathcal{A}_{bs}^{\text{C}} + \mathcal{A}_{bs}^{\text{C}} , \qquad (1.1)$$

with

$$\mathcal{A}_{b,s}^{C} = \frac{e_{p}^{2} \kappa_{b}^{2}}{4\pi} \left( \frac{\beta_{b} m_{b}}{2\pi} \right)^{1/2} v_{p} \int_{0}^{1} du \, u^{1/2} \, \exp\left\{ -\frac{1}{2} \beta_{b} m_{b} v_{p}^{2} \, u \right\}$$

$$\left[ -\ln\left( \beta_{b} \frac{e_{p} e_{b}}{4\pi} \, K \, \frac{m_{b}}{m_{pb}} \, \frac{u}{1-u} \right) - 2\gamma + 2 \right]$$
(1.2)

$$\mathcal{A}_{b,R}^{<} = \frac{e_p^2}{4\pi} \frac{i}{2\pi} \int_{-1}^{1} d\cos\theta \cos\theta \frac{\rho_b(v_p \cos\theta)}{\rho_{\text{total}}(v_p \cos\theta)} F(v_p \cos\theta) \ln\left\{\frac{F(v_p \cos\theta)}{K^2}\right\}, \quad (1.3)$$

$$\mathcal{A}_{b}^{\Delta Q} = -\frac{e_{p}^{2} \kappa_{b}^{2}}{4\pi} \left(\frac{\beta_{b} m_{b}}{2\pi}\right)^{1/2} \frac{1}{2} \int_{0}^{\infty} dv_{pb} \left\{ 2 \operatorname{Re} \psi \left(1 + i \eta_{pb}\right) - \ln \eta_{pb}^{2} \right\} 
\frac{1}{\beta_{b} m_{b} v_{p} v_{pb}} \left[ \exp \left\{ -\frac{1}{2} \beta_{b} m_{b} \left(v_{p} - v_{pb}\right)^{2} \right\} \left(1 - \frac{1}{\beta_{b} m_{b} v_{p} v_{pb}} \right) 
+ \exp \left\{ -\frac{1}{2} \beta_{b} m_{b} \left(v_{p} + v_{pb}\right)^{2} \right\} \left(1 + \frac{1}{\beta_{b} m_{b} v_{p} v_{pb}} \right) \right],$$
(1.4)

and  $\eta_{ab} = e_a e_b / 4\pi \hbar v_{ab}$ . Figures 1–3 show the  $\mathcal{A}$ -coefficients for (i)  $T_e = T_{\rm I} = 10 \,\mathrm{keV}$ , (ii)  $T_e = 10 \,\mathrm{keV}$  and  $T_{\rm I} = 100 \,\mathrm{keV}$ , and (iii)  $T_e = 100 \,\mathrm{keV}$  and  $T_{\rm I} = 10 \,\mathrm{keV}$  respectively, all with an electron number density  $n_e = 10^{25} \,\mathrm{cm}^{-3}$ .

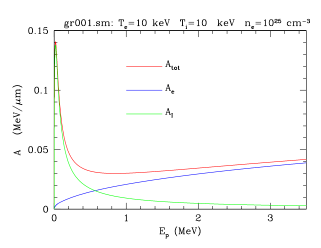


FIG. 1: Electron and ion components:  $n_e = 10^{25} \,\mathrm{cm}^{-3}$ ,  $T_e = 10 \,\mathrm{keV}$ ,  $T_{\rm I} = 10 \,\mathrm{keV}$ . [gr001.f90, gr001.sm, gr001.dat, gr001.eps]

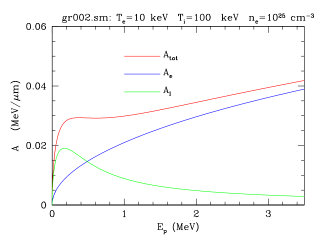


FIG. 2: Electron and ion components:  $n_e=10^{25}\,\mathrm{cm}^{-3},\ T_e=10\,\mathrm{keV},\ T_\mathrm{I}=100\,\mathrm{keV}.$  [gr002.f90, gr002.sm, gr002.dat, gr002.eps]

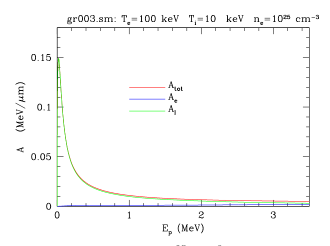


FIG. 3: Electron and ion components:  $n_e = 10^{25} \, \mathrm{cm}^{-3}$ ,  $T_e = 100 \, \mathrm{keV}$ ,  $T_{\mathrm{I}} = 10 \, \mathrm{keV}$ . [gr003.f90, gr003.sm, gr003.dat, gr003.eps]

#### II. SMALL ENERGY ASYMPTOTIC BEHAVIOR

In the small energy limit, the A-coefficients become:

$$v_p \to 0 : \mathcal{A}_b(v_p) = \underbrace{\frac{e_p^2 \kappa_b^2}{4\pi}}_{c_1} \underbrace{\left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} v_p}_{c_2} \cdot \left\{ A_b^{\text{C}} + A_b^{\Delta Q} \right\} ,$$
 (2.1)

with

$$A_b^{\rm C} = \frac{2}{3} \left[ \ln \left( \frac{16\pi}{e_p e_b \, \beta_b \kappa_{\rm D}} \frac{m_{pb}}{m_b} \right) - \frac{1}{2} - 2\gamma \right]$$
 (2.2)

$$A_b^{\Delta Q} = -\bar{\eta}_{pb}^2 \int_0^\infty du \, u \, \exp\left\{-\frac{3}{2}\bar{\eta}_{pb}^2 \, u^2\right\} \left[2 \operatorname{Re} \psi \left(1 + \frac{i}{u}\right) + \ln u^2\right] \,, \tag{2.3}$$

and  $\bar{\eta}_{pb} = e_p e_b / 4\pi \hbar \bar{v}_b$  (note:  $\bar{v}_b^2 = 3T_b / m_b$ ). The small energy quantum contribution takes separate forms for electrons and ions:

$$\bar{\eta}_{pe}^2 \ll 1 : A_e^{\Delta Q} \simeq -\bar{\eta}_{pe}^2 \int_0^\infty du \, u \, \exp\left\{-\frac{3}{2}\bar{\eta}_{pe}^2 \, u^2\right\} \left[-2\,\gamma + \ln u^2\right] = \frac{1}{3} \ln\left(\frac{3}{2}\,\bar{\eta}_{pe}^2\right) + \gamma \,.$$
(2.4)

$$\bar{\eta}_{pi}^2 \gg 1 : A_i^{\Delta Q} \simeq -\frac{\bar{\eta}_{pi}^2}{6} \int_0^\infty du \, u^3 \exp\left\{-\frac{3}{2}\bar{\eta}_{pi}^2 \, u^2\right\} = -\frac{1}{27}\,\bar{\eta}_{pi}^{-2} \,.$$
 (2.5)

The next figure illustrates the relative size of the classical contribution  $\mathcal{A}^{\text{C}}$  and the total coefficient  $\mathcal{A} = \mathcal{A}^{\text{C}} + \mathcal{A}^{\text{QM}}$ . for the density  $n_e = 10^{25} \, \text{cm}^{-3}$ . This gives us an idea of the size of the quantum contribution.

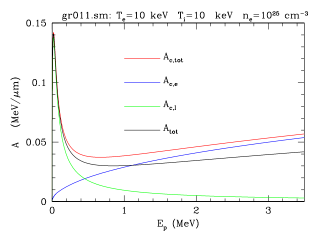


FIG. 4: The classical contributions to the A-coefficient, with the total (classical + quantum) in black. [gr001.f90, gr011.sm, gr001.dat, gr011.eps]

### A. Ions: Classical and Quantum

For each of the three cases illustrated in Figs. 1–3, we will now look at the ion contributions for the classical and quantum cases.

### 1. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_{\mathrm{I}} = 10 \,\mathrm{keV}$

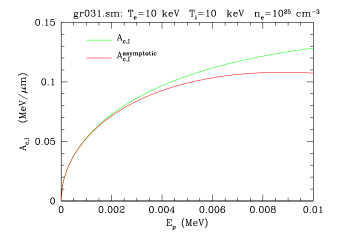


FIG. 5: Asymptotic classical ion contribution at low energies. [gr001.f90, gr031.sm, gr001.dat, gr001.smallE.dat, gr031.eps]

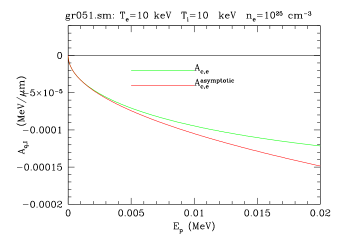


FIG. 6: Asymptotic quantum ion contribution at low energies. [gr001.f90, gr051.sm, gr001.dat, gr001.smallE.dat, gr051.eps]

# 2. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_I = 100 \,\mathrm{keV}$

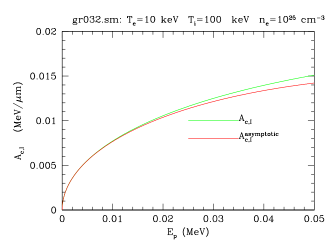


FIG. 7: Asymptotic classical ion contribution at low energies. [gr002.f90, gr032.sm, gr002.dat, gr002.smallE.dat, gr032.eps]

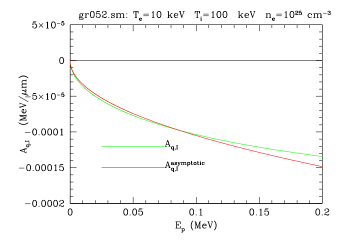


FIG. 8: Asymptotic quantum ion contribution at low energies. [gr002.f90, gr052.sm, gr002.dat, gr002.smallE.dat, gr052.eps]

# 3. Temperatures $T_e = 100 \,\mathrm{keV}$ and $T_I = 10 \,\mathrm{keV}$

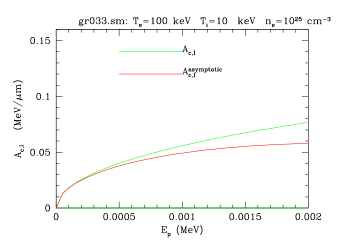


FIG. 9: Asymptotic classical ion contribution at low energies. [gr003.f90, gr033.sm, gr003.dat, gr003.smallE.dat, gr033.eps]

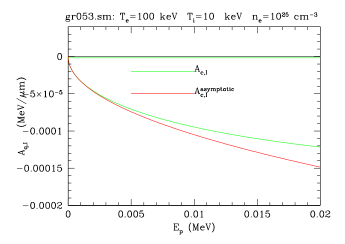


FIG. 10: Asymptotic quantum ion contribution at low energies. [gr003.f90, gr053.sm, gr003.dat, gr003.smallE.dat, gr053.eps]

### B. Electrons: Classical and Quantum

For each of the three cases illustrated in Figs. 1–3, we will now look at the electron contributions for the classical and quantum cases.

### 1. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_{\mathrm{I}} = 10 \,\mathrm{keV}$

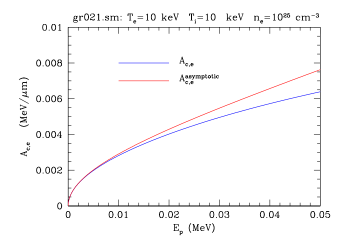


FIG. 11: Asymptotic classical electron contribution at low energies. [gr001.f90, gr021.sm, gr001.dat, gr001.smallE.dat, gr021.eps]

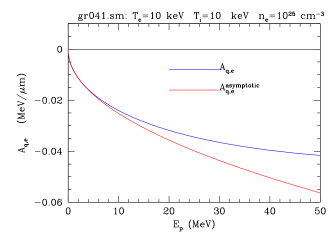


FIG. 12: Asymptotic quantum electron contribution at low energies. [gr001.f90, gr041.sm, gr001.dat, gr001.smallE.dat, gr041.eps]

# 2. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_I = 100 \,\mathrm{keV}$

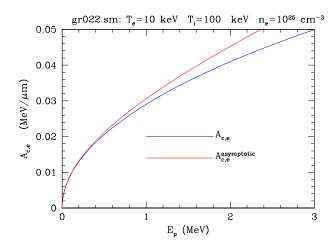


FIG. 13: Asymptotic classical electron contribution at low energies. [gr002.f90, gr022.sm, gr002.dat, gr002.smallE.dat, gr022.eps]

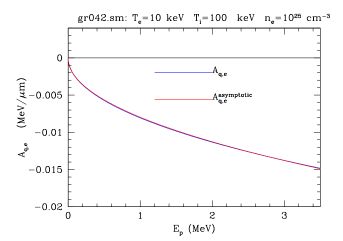


FIG. 14: Asymptotic quantum electron contribution at low energies. [gr002.f90, gr042.sm, gr002.dat, gr002.smallE.dat, gr042.eps]

# 3. Temperatures $T_e = 100 \,\mathrm{keV}$ and $T_I = 10 \,\mathrm{keV}$

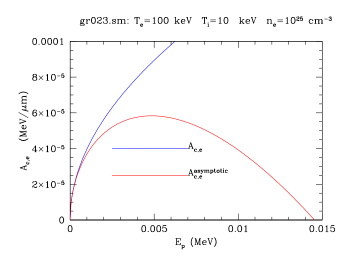


FIG. 15: Asymptotic classical electron contribution at low energies. [gr003.f90, gr023.sm, gr003.dat, gr003.smallE.dat, gr023.eps]

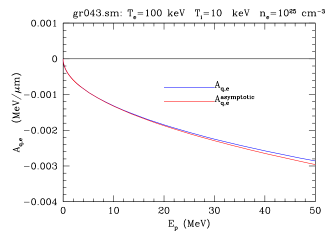


FIG. 16: Asymptotic quantum electron contribution at low energies. [gr003.f90, gr043.sm, gr003.dat, gr003.smallE.dat, gr043.eps]

### C. Total Electron and Ion Contributions

1. Temperatures  $T_e = 10 \,\mathrm{keV}$  and  $T_{\mathrm{I}} = 10 \,\mathrm{keV}$ 

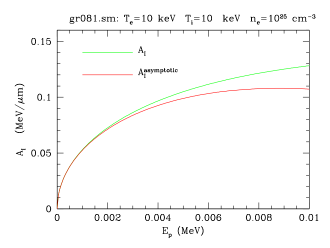


FIG. 17: Total asymptotic ion contribution at low energies. [gr001.f90, gr081.sm, gr001.dat, gr001.smallE.dat, gr081.eps]

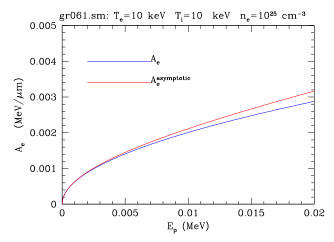


FIG. 18: Total asymptotic electron contribution at low energies. [gr001.f90, gr061.sm, gr001.dat, gr001.smallE.dat, gr061.eps]

# 2. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_I = 100 \,\mathrm{keV}$

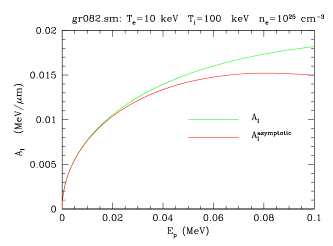


FIG. 19: Total asymptotic ion contribution at low energies. [gr002.f90, gr082.sm, gr002.dat, gr002.smallE.dat, gr082.eps]

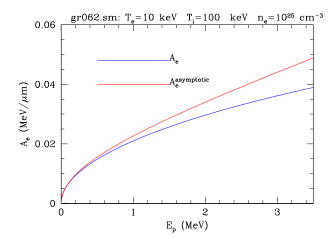


FIG. 20: Total asymptotic electron contribution at low energies. [gr002.f90, gr062.sm, gr002.dat, gr002.smallE.dat, gr062.eps]

# 3. Temperatures $T_e = 100 \,\mathrm{keV}$ and $T_I = 10 \,\mathrm{keV}$

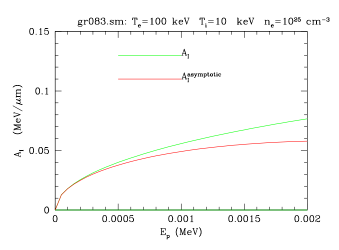


FIG. 21: Total asymptotic ion contribution at low energies. [gr003.f90, gr083.sm, gr003.dat, gr003.smallE.dat, gr083.eps]

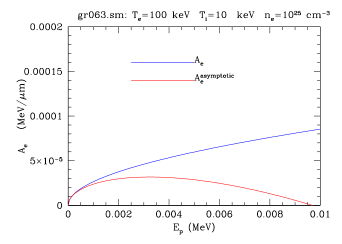


FIG. 22: Total asymptotic electron contribution at low energies. [gr003.f90, gr063.sm, gr003.dat, gr003.smallE.dat, gr063.eps]

#### D. More Classical: The Singular and Regular Contributions

Up to now, we have not been dividing the classical piece into its singular and regular contributions.

$$v_p \to 0 : \mathcal{A}_b(v_p) = \underbrace{\frac{e_p^2 \kappa_b^2}{4\pi}}_{c_1} \underbrace{\left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} v_p}_{c_2} \cdot \left\{ \left(\bar{A}_{b,s}^{\text{CL}} + \bar{A}_{b,R}^{<}\right) + A_b^{\Delta Q} \right\},$$
 (2.6)

Since the code calculates the singular and regular pieces separately, let's explore their respective small energy asymptotics. These are given by Eq. (9.8) on p. 300 and Eq. (7.34) on p (287) of BPS, respectively:

$$\bar{A}_{b,s}^{CL} = -\left(\frac{2}{3} - \frac{1}{5}\beta_b m_b v_p^2\right) \left[\ln\left\{\frac{e_p e_b \beta_b K}{16\pi} \frac{m_b}{m_{pb}}\right\} + 2\gamma\right] + \frac{2}{15}\beta_b m_b v_p^2 + \mathcal{O}(v_p^4) ; \quad (2.7)$$

$$\bar{A}_{b,R}^{<} = -\left(\frac{2}{3} - \frac{1}{5}\beta_b m_b v_p^2\right) \left[\frac{1}{2} + \ln\left\{\frac{\kappa_D}{K}\right\}\right] + \frac{1}{5} \sum_c \frac{\kappa_c^2}{\kappa_D^2} \beta_c m_c v_p^2 - \frac{\pi}{36} \left[\sum_c \frac{\kappa_c^2}{\kappa_D^2} \left(\beta_c m_c v_p^2\right)^{1/2}\right]^2 + \mathcal{O}(v_p^4) . \tag{2.8}$$

(2.9)

To leading order in  $v_p$  these expressions become

$$\bar{A}_{b,s}^{CL} = -\frac{2}{3} \left[ \ln \left\{ \frac{e_p \, e_b \beta_b \, K}{16\pi} \, \frac{m_b}{m_{pb}} \right\} + 2\gamma \right] + \mathcal{O}(v_p^2)$$
 (2.10)

$$\bar{A}_{b,R}^{<} = -\frac{2}{3} \left[ \frac{1}{2} + \ln \left\{ \frac{\kappa_{\rm D}}{K} \right\} \right] + \mathcal{O}(v_p^2) ,$$
 (2.11)

and upon adding them the K's cancel and we return to (2.2). Keep the  $v_p^2$  terms, we have the more precise expression (this is the one actually used in the code):

$$\bar{A}_{b,s}^{CL} = -\left(\frac{2}{3} - \frac{1}{5}\beta_b m_b v_p^2\right) \left[\ln\left\{\frac{e_p e_b \beta_b K}{16\pi} \frac{m_b}{m_{pb}}\right\} + 2\gamma\right] + \frac{2}{15}\beta_b m_b v_p^2 + \mathcal{O}(v_p^4) \quad (2.12)$$

$$\bar{A}_{b,R}^{<} = -\left(\frac{2}{3} - \frac{1}{5}\beta_b m_b v_p^2\right) \left[\frac{1}{2} + \ln\left\{\frac{\kappa_D}{K}\right\}\right] + \frac{1}{5} \sum_c \frac{\kappa_c^2}{\kappa_D^2} \beta_c m_c v_p^2 - \frac{\pi}{36} \left[\sum_c \frac{\kappa_c^2}{\kappa_D^2} \left(\beta_c m_c v_p^2\right)^{1/2}\right]^2 + \mathcal{O}(v_p^4) , \qquad (2.13)$$

$$\bar{A}_{b}^{\text{CL}} = \bar{A}_{b,\text{S}}^{\text{CL}} + \bar{A}_{b,\text{R}}^{<} = -\left(\frac{2}{3} - \frac{1}{5}\beta_{b}m_{b}v_{p}^{2}\right) \left[\ln\left\{\frac{e_{p}e_{b}\beta_{b}\kappa_{\text{D}}}{16\pi} \frac{m_{b}}{m_{pb}}\right\} + \frac{1}{2} + 2\gamma\right] + \frac{2}{15}\beta_{b}m_{b}v_{p}^{2}$$

$$\frac{\pi}{36} \left[\sum_{c} \frac{\kappa_{c}^{2}}{\kappa_{\text{D}}^{2}} \left(\beta_{c}m_{c}v_{p}^{2}\right)^{1/2}\right]^{2} + \mathcal{O}(v_{p}^{4}) .$$
(2.14)

# 1. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_I = 10 \,\mathrm{keV}$

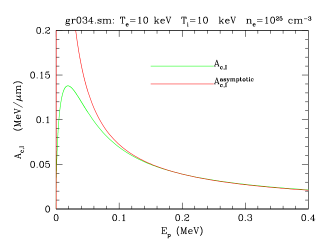


FIG. 23: Asymptotic singular ion contribution at low energies. [gr004.f90, gr004.dat, gr004.smallE.dat, gr104.sm]

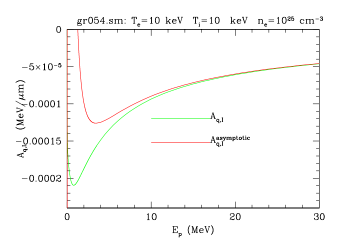


FIG. 24: Asymptotic regular ion contribution at low energies. \*\*\* [gr001.f90, gr034.sm, gr001.dat, gr001.highE.dat, gr034.eps] \*\*\*

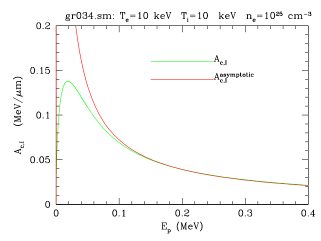


FIG. 25: Asymptotic singular electron contribution at low energies. \*\*\* [gr001.f90, gr034.sm, gr001.dat, gr001.highE.dat, gr034.eps] \*\*\*

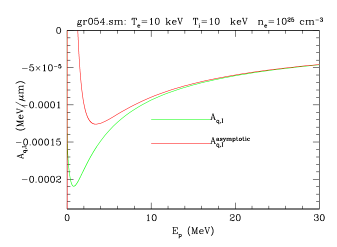


FIG. 26: Asymptotic regular electron contribution at low energies. \*\*\* [gr001.f90, gr034.sm, gr001.dat, gr001.highE.dat, gr034.eps] \*\*\*

#### III. HIGH ENERGY ASYMPTOTIC BEHAVIOR

#### A. Ions: Classical and Quantum

We will look at ion contributions before looking at the electrons (this is because there are two high energy electron regimes). For high energy  $E \gg T$ , the ion contribution to the  $\mathcal{A}$ -coefficient becomes

$$E_p \gg T : \mathcal{A}_{\text{\tiny I}}^{\text{\tiny C}} = \frac{e_p^2}{4\pi} \frac{1}{v_p^2} \sum_i \omega_i^2 \left[ -\ln \left\{ \frac{e_p e_i \kappa_e}{16\pi} \frac{2}{m_{pi} v_p^2} \right\} - \gamma - \frac{1}{2} \right]$$
 [(B36) in text] (3.1)

$$\mathcal{A}_{I}^{QM} = \frac{e_{p}^{2}}{4\pi} \frac{1}{v_{p}^{2}} \sum_{i} \omega_{i}^{2} \left[ -\ln \left\{ \frac{\hbar \kappa_{e}}{2m_{pi}v_{p}} \right\} - \frac{1}{2} \right] . \tag{3.2}$$

To see this, note that the classical singular ion contribution at high energy is given by (B40) [v3.6]:

$$E_p \gg T \quad \eta_{pi} = \frac{e_p e_i}{4\pi \hbar v_p} \gg 1:$$

$$\mathcal{A}_{i,s}^{C} = -\frac{e_p^2}{4\pi} \frac{\omega_i^2}{v_p^2} \left[ \ln \left\{ \frac{e_p e_i}{16\pi} \frac{2\kappa_e}{m_{pi} v_p^2} \right\} + \gamma \right] \quad [(B40) \text{ in text}]$$
(3.3)

The regular high energy asymptotic ion term is

$$E_p \gg T \ln \left\{ \frac{m_{\rm I} T_e^3}{m_e T_{\rm I}^3} \right\}$$

$$\mathcal{A}_{i,\rm R}^{\rm C} = -\frac{e_p^2}{4\pi} \frac{\omega_i^2}{2v_p^2} \quad [(\rm B34) \text{ in text w/o sum}] . \tag{3.4}$$

The asymptotic quantum piece is given by subtracting (B36) for the singular contribution from (B42) [the singular + quantum], to give

$$E_{p} \gg T: \quad \eta_{pi} = \frac{e_{p}e_{i}}{4\pi\hbar v_{p}}$$

$$\mathcal{A}_{i}^{QM} = \frac{e_{p}^{2}}{4\pi} \frac{\omega_{i}^{2}}{v_{p}^{2}} \left[ \ln \eta_{pi} + \gamma \right] \quad \text{note:} \quad \frac{e_{p}^{2}}{4\pi} \frac{\omega_{i}^{2}}{v_{p}^{2}} = \frac{e_{p}^{2} \kappa_{i}^{2}}{4\pi} \frac{\omega_{i}^{2}}{\kappa_{i}^{2} v_{p}^{2}} = \frac{e_{p}^{2} \kappa_{i}^{2}}{4\pi} \frac{1}{m_{i} \beta_{i} v_{p}^{2}}. \quad (3.5)$$

The algebra is:

$$\mathcal{A}_{i}^{\text{QM}} = (B42) - (B36)$$

$$= e_{p}^{2} \omega_{i}^{2} \left[ \ln \int 2m_{pi}v_{p} \right] \qquad e_{p}^{2} \omega_{i}^{2} \left[ \ln \int e_{p}e_{i}\kappa_{e} \quad 2 \right]$$

$$(3.6)$$

$$= \frac{e_p^2}{4\pi} \frac{\omega_i^2}{v_p^2} \left[ \ln \left\{ \frac{2m_{pi}v_p}{\hbar \kappa_e} \right\} \right] - \frac{e_p^2}{4\pi} \frac{\omega_i^2}{v_p^2} \left[ -\ln \left\{ \frac{e_p e_i \kappa_e}{16\pi} \frac{2}{m_{pi} v_p^2} \right\} - \gamma \right]$$
(3.7)

$$= \frac{e_p^2}{4\pi} \frac{\omega_i^2}{v_p^2} \left[ \ln \left\{ \frac{2m_{pi}v_p}{\hbar\kappa_e} \cdot \frac{e_p e_i \kappa_e}{16\pi} \frac{2}{m_{pi}v_p^2} \right\} + \gamma \right]$$
 (3.8)

$$= \frac{e_p^2}{4\pi} \frac{\omega_i^2}{v_p^2} \left[ \ln \left\{ \frac{e_p e_i}{4\pi \, \hbar v_p} \right\} + \gamma \right] . \tag{3.9}$$

# 1. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_I = 10 \,\mathrm{keV}$

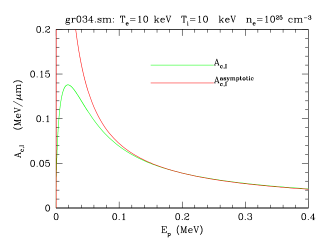


FIG. 27: Asymptotic classical ion contribution at high energies. [gr001.f90, gr034.sm, gr001.dat, gr001.high E.dat, gr034.eps]

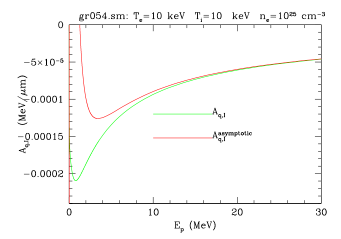


FIG. 28: Asymptotic quantum ion contribution at high energies. [gr001.f90, gr054.sm, gr001.dat, gr001.high E.dat, gr054.eps]

# 2. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_I = 100 \,\mathrm{keV}$

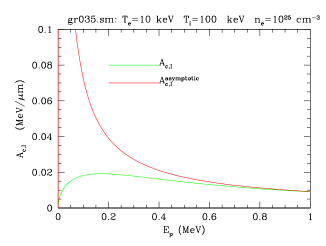


FIG. 29: Asymptotic classical ion contribution at high energies. [gr002.f90, gr035.sm, gr002.dat, gr002.high E.dat, gr035.eps]

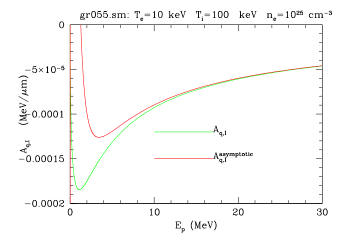


FIG. 30: Asymptotic quantum ion contribution at high energies. [gr002.f90, gr055.sm, gr002.dat, gr002.high E.dat, gr055.eps]

# 3. Temperatures $T_e = 100 \,\mathrm{keV}$ and $T_I = 10 \,\mathrm{keV}$

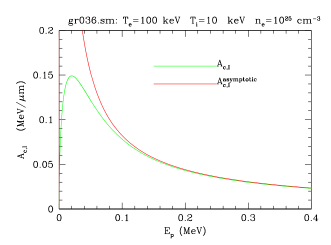


FIG. 31: Asymptotic classical ion contribution at high energies. [gr003.f90, gr036.sm, gr003.dat, gr003.high E.dat, gr036.eps]

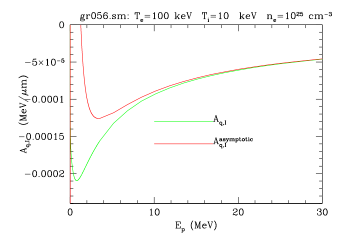


FIG. 32: Asymptotic quantum ion contribution at high energies. [gr003.f90, gr056.sm, gr003.dat, gr003.high E.dat, gr056.eps]

#### B. Electrons: Classical and Quantum

The high energy electron regime is broken into two widely separated scales, one given by T and the other by  $m_1T/m_e$ . In the high-intermediate energy regime,

$$\frac{m_{\rm I}}{m_e}\,T \gg E_p \gg T: \quad {\rm DT~plasma~at~10\,keV} \Rightarrow \frac{m_{\rm I,\,av}}{m_e}\,T = \frac{2.5\,{\rm GeV}}{10\,{\rm keV}} = 2.5\,{\rm MeV}.$$

$$\mathcal{A}_{e,R}^{<} = -\frac{e_p^2 \kappa_e^2}{4\pi} \left(\frac{\beta_e m_e}{2\pi}\right)^{1/2} \frac{v_p}{3} \left[ \ln \left\{ \frac{\kappa_e^2}{K^2} \right\} + 1 \right] \quad [\text{ (B51) in text}]$$
 (3.10)

$$\mathcal{A}_{e, s}^{C} + \mathcal{A}_{e}^{QM} = \frac{e_{p}^{2} \kappa_{e}^{2}}{4\pi} \left(\frac{\beta_{e} m_{e}}{2\pi}\right)^{1/2} \frac{v_{p}}{3} \left[ \ln \left\{ \frac{8T_{e} m_{pe}^{2}}{m_{e} \hbar^{2} K^{2}} \right\} - \gamma \right] \quad [\text{ (B55) in text}]$$
(3.11)

$$\mathcal{A}_{e} = \mathcal{A}_{e,R}^{<} + \mathcal{A}_{e,S}^{C} + \mathcal{A}_{e}^{QM} 
= \frac{e_{p}^{2} \kappa_{e}^{2}}{4\pi} \left(\frac{\beta_{e} m_{e}}{2\pi}\right)^{1/2} \frac{v_{p}}{3} \left[ \ln \left\{ \frac{8T_{e} m_{pe}^{2}}{m_{e} \hbar^{2} \kappa_{e}^{2}} \right\} - \gamma - 1 \right] \quad [\text{ (B56) in text]}, \quad (3.12)$$

and in the extreme high energy regime,

$$E_p \gg \frac{m_{\rm I}}{m_e} T:$$

$$\mathcal{A}_e = \frac{e_p^2}{4\pi} \frac{\omega_e^2}{v^2} \ln \left\{ \frac{2m_{pe}v_p^2}{\hbar\omega_e} \right\} \quad [\text{ (B58) in text]}. \tag{3.13}$$

We now have the total electron contribution  $\mathcal{A}_e$  in the extreme and intermediate high energy regimes; however, I will have to calculate  $\mathcal{A}_e^{\text{QM}}$  later since the text does not give  $\mathcal{A}_{e, s}^{\text{C}}$  (as it did for the ions).

I will use the coding notation:

$$a_b = \frac{1}{2} \beta_b m_b v_p^2 = \frac{1}{2} \beta_b m_b c^2 \left( v_p^2 / c^2 \right)$$
 (3.14)

$$c_2 = \left(\frac{a_e}{\pi}\right)^{1/2} = \left(\frac{\beta_e m_e}{2\pi}\right)^{1/2} v_p = \left(\frac{\beta_e m_e c^2}{2\pi}\right)^{1/2} \frac{v_p}{c}$$
 (3.15)

$$c_1 = \frac{e_p^2 \kappa_e^2}{4\pi} = Z_p^2 \cdot \frac{e^2}{8\pi a_0} \cdot 2a_0 \cdot \kappa_e^2 = 2Z_p^2 B_e \kappa_e^2 a_0 . \tag{3.16}$$

For coding purposes, the extreme high energy electron contribution can then be written,

$$\mathcal{A}_{e} = c_{1} \frac{\omega_{e}^{2}}{\kappa_{e}^{2} v_{p}^{2}} \ln \left\{ \frac{2 (m_{pe}c^{2}) v_{p}^{2}}{(\hbar c) c \omega_{e}} \right\} = \frac{c_{1}}{\beta_{e} m_{e} v_{p}^{2}} \ln \left\{ \frac{2 (m_{pe}c^{2}) v_{p}^{2}}{(\hbar c) c \omega_{e}} \right\}$$
(3.17)

$$= \frac{c_1}{2a_e} \ln \left\{ \frac{2(m_{pe}c^2)v_p^2}{(\hbar c) c \omega_e} \right\}$$
 (B58), (3.18)

and the intermediate high energy form is

$$\mathcal{A}_{e} = \frac{c_{1} c_{2}}{3} \left[ \ln \left\{ \frac{8T_{e} (m_{pe}c^{2})^{2}}{(m_{e}c^{2})(\hbar c)^{2} \kappa_{e}^{2}} \right\} - \gamma - 1 \right]$$
 (B56) . (3.19)

1. Temperatures  $T_e = 10 \,\mathrm{keV}$  and  $T_I = 10 \,\mathrm{keV}$ 

FIG. 33: Asymptotic classical electron contribution at high energies. [gr001.f90, gr024.sm, gr001.dat, gr001.highE.dat, gr024.eps]

FIG. 34: Asymptotic quantum electron contribution at high energies. [gr001.f90, gr044.sm, gr001.dat, gr001.highE.dat, gr044.eps]

2. Temperatures  $T_e = 10 \,\mathrm{keV}$  and  $T_I = 100 \,\mathrm{keV}$ 

FIG. 35: Asymptotic classical electron contribution at high energies. [gr002.f90, gr025.sm, gr002.dat, gr002.highE.dat, gr025.eps]

FIG. 36: Asymptotic quantum electron contribution at high energies. [gr002.f90, gr045.sm, gr002.dat, gr002.highE.dat, gr045.eps]

3. Temperatures  $T_e = 100 \,\mathrm{keV}$  and  $T_I = 10 \,\mathrm{keV}$ 

FIG. 37: Asymptotic classical electron contribution at high energies. [gr003.f90, gr026.sm, gr003.dat, gr003.highE.dat, gr026.eps]

FIG. 38: Asymptotic quantum electron contribution at high energies. [gr003.f90, gr046.sm, gr003.dat, gr003.highE.dat, gr046.eps]

### C. Total Electron and Ion Contributions

# 1. Temperatures $T_e=10\,\mathrm{keV}$ and $T_\mathrm{I}=10\,\mathrm{keV}$

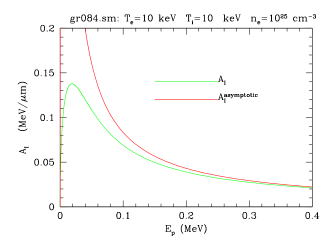


FIG. 39: Total asymptotic ion contribution at high energies. [gr001.f90, gr084.sm, gr001.dat, gr001.highE.dat, gr084.eps]

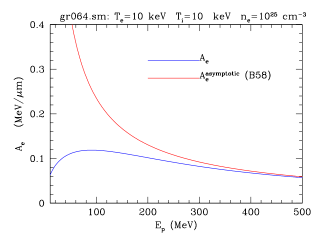


FIG. 40: Total asymptotic electron contribution (B58) at very high energies. [gr001.f90, gr064.sm, gr001.dat, gr001.very.highE.dat, gr064.eps]

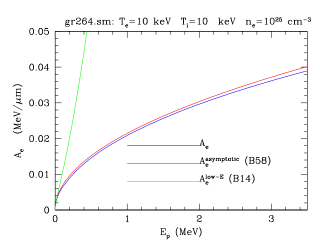


FIG. 41: Total asymptotic electron contribution (B56) at medium high energies. [gr001.f90, gr264.sm, gr001.dat, gr001.highE.dat, gr264.eps]

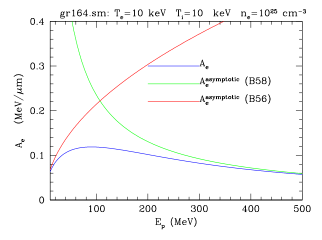


FIG. 42: Total asymptotic electron contribution (B56) and (B58). [gr001.f90, gr164.sm, gr001.dat, gr001.highE.dat, gr001.very.highE.dat, gr164.eps]

# 2. Temperatures $T_e = 10 \,\mathrm{keV}$ and $T_I = 100 \,\mathrm{keV}$

 $gr085.sm: \ T_{e}{=}10 \ keV \ T_{i}{=}100 \ keV \ n_{e}{=}10^{25} \ cm^{-3}$ 

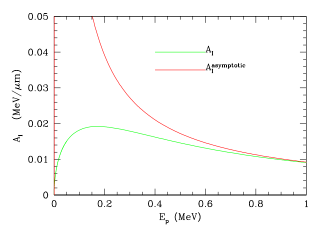


FIG. 43: Total asymptotic ion contribution at high energies. [gr002.f90, gr085.sm, gr002.dat, gr002.high E.dat, gr085.eps]

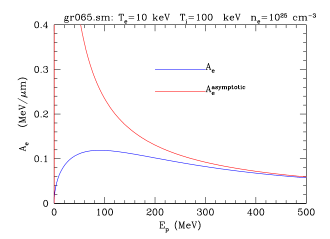


FIG. 44: Total asymptotic electron contribution (B58) at very high energies. [gr002.f90, gr065.sm, gr002.dat, gr002.highE.dat, gr065.eps]

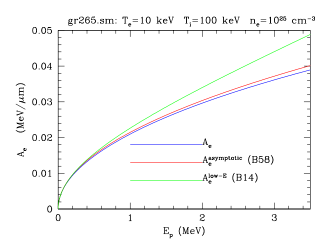


FIG. 45: Total asymptotic electron contribution (B56) at medium high energies. [gr002.f90, gr265.sm, gr002.dat, gr002.highE.dat, gr265.eps]

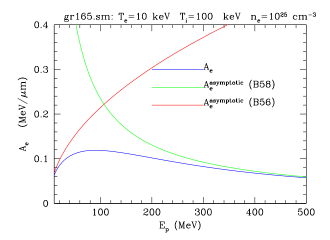


FIG. 46: Total asymptotic electron contribution (B56) and (B58). [gr002.f90, gr165.sm, gr002.dat, gr002.highE.dat, gr002.very.highE.dat, gr165.eps]

# 3. Temperatures $T_e = 100 \,\mathrm{keV}$ and $T_I = 10 \,\mathrm{keV}$

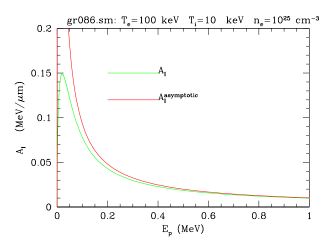


FIG. 47: Total asymptotic ion contribution at high energies. [gr003.f90, gr086.sm, gr003.dat, gr003.high E.dat, gr086.eps]

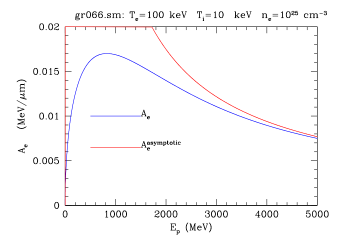


FIG. 48: Total asymptotic electron contribution (B58) at very high energies. [gr003.f90, gr066.sm, gr003.dat, gr003.very.highE.dat, gr066.eps]

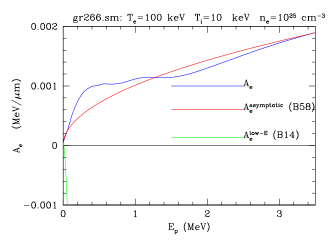


FIG. 49: Total asymptotic electron contribution (B56) at medium high energies. [gr003.f90, gr266.sm, gr003.dat, gr003.highE.dat, gr266.eps]

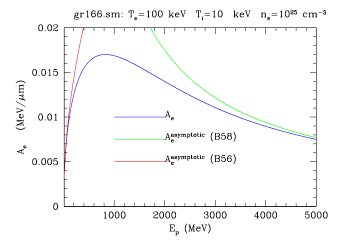


FIG. 50: Total asymptotic electron contribution (B56) at medium high energies. [gr003.f90, gr166.sm, gr003.dat, gr003.highE.dat, gr003.very.highE.dat, gr166.eps]

### Appendix A: Coding the A-coefficients

#### 1. The Singular Contribution

The singular contribution,

$$\mathcal{A}_{b,s}^{C} = \left[ \frac{e_{p}^{2} \kappa_{b}^{2}}{4\pi} \left( \frac{\beta_{b} m_{b}}{2\pi} \right)^{1/2} v_{p} \right] \int_{0}^{1} du \, u^{1/2} e^{-\frac{1}{2} \beta_{b} m_{b} v_{p}^{2} u} \left[ -\ln \left\{ \frac{\beta_{b} e_{b} e_{p}}{4\pi} K \frac{m_{b}}{m_{pb}} \frac{u}{1-u} \right\} - 2\gamma + 2 \right], \tag{A1}$$

is quite easy to code. The integral can be broke into the pieces

$$\int_{0}^{1} du \, u^{1/2} e^{-\frac{1}{2}\beta_{b} m_{b} v_{p}^{2} u} \left[ \ln \left\{ \frac{u}{1-u} \right\} - \ln \left\{ \frac{\beta_{b} e_{b} e_{p}}{4\pi} K \frac{m_{b}}{m_{pb}} \right\} - 2\gamma + 2 \right], \tag{A2}$$

which motivates the definition

$$\mathcal{A}_{b,S}^{C} = c_{b,1} c_{b,2} \cdot \mathsf{A}_{S}(a_{pb}, b_{pb}) \tag{A3}$$

$$A_{s}(a,b) = \int_{0}^{1} du \, u^{1/2} e^{-a \, u} \left[ -\ln \left\{ \frac{u}{1-u} \right\} + b \right]$$
(A4)

$$a_{pb} = \frac{1}{2} \beta_b m_b v_p^2$$
 and  $b_{pb} = -\ln \left\{ \frac{\beta_b e_b e_p}{4\pi} K \frac{m_b}{m_{pb}} \right\} - 2\gamma + 2$  (A5)

$$c_{b,1} = \frac{e_p^2 \kappa_b^2}{4\pi} \quad c_{b,2} = \left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} v_p .$$
 (A6)

The term involving b can be integrated exactly, but we will use Gaussian quadrature for both pieces.

acoeff.f90:

The numerical integration is performed by Gaussian quadrature:

acoeff.f90:

```
SUBROUTINE a_sing(a, b, ac_s)

REAL, INTENT(IN) :: a

REAL, INTENT(IN) :: b

REAL, INTENT(OUT) :: ac_s

REAL :: u0, u1, du, um

INTEGER, PARAMETER :: NS=1000 ! integration regions singular: must

REAL, PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad

REAL, PARAMETER :: W13=0.5555555556E0, W2=0.8888888889E0

ac_s=0
```

```
u0=0
u1=1
du=(u1-u0)/NS
u=u0-du
D0 iu=1,NS,2 ! Gaussian quadrature
u=u+2.E0*du
ac_s=ac_s+W2*dab_sing(u,a,b)
um=u-du*UPM
ac_s=ac_s+W13*dab_sing(um,a,b)
um=u+du*UPM
ac_s=ac_s+W13*dab_sing(um,a,b)
ENDD0
ac_s=ac_s*du
END SUBROUTINE a_sing
```

#### 2. The Regular Contribution

The long-distance regular contribution can be expressed as

$$\mathcal{A}_{b,R}^{<} = \frac{e_p^2}{4\pi} \frac{i}{2\pi} \int_{-1}^{1} du \, u \, \frac{\rho_b(v_p u)}{\rho_{\text{total}}(v_p u)} F(v_p u) \ln\left\{\frac{F(v_p u)}{K^2}\right\} \tag{A7}$$

$$= \frac{e_p^2}{4\pi} \frac{i}{2\pi} \int_0^1 du \, u \, \frac{\rho_b(v_p u)}{\rho_{\text{total}}(v_p u)} \left[ F(v_p u) \ln \left\{ \frac{F(v_p u)}{K^2} \right\} - F^*(v_p u) \ln \left\{ \frac{F^*(v_p u)}{K^2} \right\} \right] (A8)$$

$$= -\frac{e_p^2}{4\pi} \frac{1}{2\pi} \int_0^1 du \, u \, \frac{\rho_b(v_p u)}{\rho_{\text{total}}(v_p u)} \, H(v_p u) \,, \tag{A9}$$

where we have defined

$$H(v) \equiv -i \left[ F(v) \ln \left\{ \frac{F(v)}{K^2} \right\} - F^*(v) \ln \left\{ \frac{F^*(v)}{K^2} \right\} \right] = 2 \left[ F_{\text{Re}} \arg\{F\} + F_{\text{Im}} \ln \left\{ \frac{|F|}{K^2} \right\} \right]. \tag{A10}$$

We shall factor out a dimensionfull wavenumber K and define dimensionless quantities  $\mathbb{F}(v)$  and  $\mathbb{H}(v)$  through

$$F(v) = K^2 \mathbb{F}(v)$$
 and  $H(v) = K^2 \mathbb{H}(v)$ . (A11)

Defining the parameters

$$a_c \equiv \left(\frac{\beta_c m_c}{2}\right)^{1/2} \tag{A12}$$

$$\bar{\kappa}_c^2 \equiv \frac{\kappa_c^2}{K^2} \tag{A13}$$

gives the real and imaginary parts of  $\mathbb{F}$ .

$$\mathbb{F}_{\text{Re}}(\{a_c v\}, \{\bar{\kappa}_c\}) = \sum_{c} \bar{\kappa}_c^2 \left(1 - 2a_c v \operatorname{daw}\{a_c v\}\right)$$
(A14)

$$\mathbb{F}_{\text{Im}}(\{a_c v\}, \{\bar{\kappa}_c\}) = \sqrt{\pi} \sum_{c} \bar{\kappa}_c^2 a_c v e^{-a_c^2 v^2}. \tag{A15}$$

The ratio of weighting factors can be written

$$\frac{\kappa_b^2}{K^2} \mathbb{R}_b(\{(u \,\beta_c m_c v_p^2/2)^{1/2}\}) \equiv \frac{\rho_b(v_p u)}{\rho_{\text{total}}(v_p u)} = \frac{\kappa_b^2 (\beta_b m_b/2\pi)^{1/2} v_p u \, e^{-\frac{1}{2} \beta_b m_b v_p^2 u^2}}{\sum_c \kappa_c^2 (\beta_c m_c/2\pi)^{1/2} v_p u \, e^{-\frac{1}{2} \beta_c m_c v_p^2 u^2}} \qquad (A16)$$

$$= \left[ \sum_c \frac{\kappa_c^2}{\kappa_b^2} \left( \frac{\beta_c m_c}{\beta_b m_b} \right)^{1/2} e^{\frac{1}{2} (\beta_b m_b - \beta_c m_c) v_p^2 u^2} \right]^{-1}, \qquad (A17)$$

or

$$\mathbb{R}_{b}(\{(u\,\beta_{c}m_{c}v_{p}^{2}/2)^{1/2}\,\}) = \left[\sum_{c} \frac{\kappa_{c}^{2}}{K^{2}} \left(\frac{\beta_{c}m_{c}}{\beta_{b}m_{b}}\right)^{1/2} e^{\frac{1}{2}(\beta_{b}m_{b} - \beta_{c}m_{c})v_{p}^{2}u^{2}}\right]^{-1} . \tag{A18}$$

We can now express the regular piece as

$$\mathcal{A}_{b,R}^{C} = \underbrace{\left[\frac{e_p^2 \kappa_b^2}{4\pi}\right]}_{c_{b,1}} \cdot \mathsf{A}_{R}(v_p, \{a_c\}, \{\bar{\kappa}_c\})$$
(A19)

$$\mathsf{A}_{b_{\mathrm{R}}}(v_{p}, \{a_{c}\}, \{\bar{\kappa}_{c}\}) = \int_{0}^{1} du \underbrace{\mathbb{R}_{b}(\{a_{c}v_{p}u\}) \mathbb{H}(\{a_{c}v_{p}u\}, \{\bar{\kappa}_{c}\})}_{\text{dab\_reg}}. \tag{A20}$$

acoeff.f90:

```
FUNCTION dab_reg(u, vp, ib, nni, k2, kb2, betab, mb)
    USE mathvars
    USE physvars
IMPLICIT NONE
      REAL,
                                       INTENT(IN)
                                                                     [dimensionless]
                                                     :: u
      REAL,
                                       INTENT(IN)
                                                     :: vp
                                                                    Projectile velocity [c
      INTEGER,
                                       INTENT(IN)
                                                     :: ib
                                                                    Species number
      INTEGER,
                                       INTENT(IN)
                                                     :: nni
                                                                 ! Number of ion species
                                                                    Wavenumber squared [1/
      REAL,
                                       INTENT(IN)
                                                     :: k2
                                                     :: kb2
                DIMENSION(1:nni+1), INTENT(IN)
      REAL,
                                                                    Debye wavenumber squar
      REAL,
                DIMENSION(1:nni+1), INTENT(IN)
                                                    :: betab
                                                                    Temperature array [1/k
      REAL,
                DIMENSION(1:nni+1), INTENT(IN)
                                                    :: mb
                                                                    Mass array [keV]
      REAL
                                                     :: dab_reg !
                                                                     [dimensionless]
      REAL,
                DIMENSION(1:nni+1) :: alfb, ab
                                      :: fr, fi, fabs, farg, h
:: kcb, r_ib, bm_ic, bm_ib, a_ic, a_ib, ex, au
      REAL
      REAL
      INTEGER
                                      :: ic
      ab=SQRT(0.5*betab*mb)*vp/CC
      alfb=kb2/k2
      CALL frfi(u,nni,alfb,ab,fr,fi,fabs,farg)
      h=2*(fr*farg + fi*LOG(fabs))
construct spectral weight ratio Rb=rho_b/rho_tot
      r_{ib=0}
      bm_ib=betab(ib)*mb(ib)
      a_{ib} = ab(ib)*ab(ib)
      DO ic=1,nni+1
          kcb=kb2(ic)/k2
          bm_ic=betab(ic)*mb(ic)
          a_ic =ab(ic)*ab(ic)
          \overline{IF} (ic == ib) THEN
             au=(a_ic-a_ib)*u
ex=EXP(-au)
      r_ib=r_ib + kcb*SQRT(bm_ic/bm_ib)*ex ENDDO
          ENDIF
      r_ib=1/r_ib
    dab_reg=-u*r_ib*h/TWOPI
END FUNCTION dab_reg
```

The numerical integration is performed by Gaussian quadrature:

acoeff.f90:

```
SUBROUTINE a_reg(ib, nni, vp, k2, kb2, betab, mb, ac_r)
INTEGER,
INTENT(IN) :: ib
```

```
INTENT(IN)
INTENT(IN)
INTENT(IN)
INTENT(IN)
DIMENSION(1:nni+1), INTENT(IN)
DIMENSION(1:nni+1), INTENT(IN)
INTENT(UT)
                                                                 :: nni
:: k2
   INTEGER, REAL,
   ŘEAL,
                                                                :: kb2
:: betab
   REAL,
   REAL,
                                                                 :: mb
   REAL,
                                                                 :: ac_r
   REAL :: u0, u1, du, u, um INTEGER, PARAMETER :: NR=10
                PARAMETER :: NR=10 ! integration regions singular: must b PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad PARAMETER :: W13=0.555555556E0, W2=0.8888888889E0
   REAL,
   REAL,
   ac_r=0
   u0=0.
   u1=1.
   du=(u1-u0)/NR
   u=u0-du
   DO iu=1,NR,2 ! Gaussian quadrature
       u=u+2.E0*du
       ac_r=ac_r+W2*dab_reg(u,vp,ib,nni,k2,kb2,betab,mb)
       um=u-du*UPM
       ac_r=ac_r+W13*dab_reg(um,vp,ib,nni,k2,kb2,betab,mb)
       um=u+du*UPM
       ac_r=ac_r+W13*dab_reg(um,vp,ib,nni,k2,kb2,betab,mb)
   ENDDO
ac_r=ac_r*du
END SUBROUTINE a_reg
```

#### 3. Quantum Contribution

For the quantum term we make the change of variables  $v_{pb} = v_p u$  so that

$$\mathcal{A}_{b}^{\text{QM}} = -\frac{e_{p}^{2} \kappa_{b}^{2}}{4\pi} \left( \frac{\beta_{b} m_{b}}{2\pi} \right)^{1/2} v_{p} \int_{0}^{\infty} du \left[ \text{Re} \, \psi \left\{ 1 + i \, \frac{\bar{\eta}_{pb}}{u} \right\} - \ln \left\{ \frac{\bar{\eta}_{pb}}{u} \right\} \right] \frac{1}{\beta_{b} m_{b} v_{p}^{2} u}$$

$$\left[ e^{-\frac{1}{2} \beta_{b} m_{b} v_{p}^{2} (u-1)^{2}} \left( 1 - \frac{1}{\beta_{b} m_{b} v_{p}^{2} u} \right) + e^{-\frac{1}{2} \beta_{b} m_{b} v_{p}^{2} (u+1)^{2}} \left( 1 + \frac{1}{\beta_{b} m_{b} v_{p}^{2} u} \right) \right] . \quad (A21)$$

The quantum function we need to code is therefore

$$\mathcal{A}_b^{\text{QM}} = \underbrace{\left[\frac{e_p^2 \kappa_b^2}{4\pi} \left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} v_p\right]}_{c_{b,1} \cdot c_{b,2}} \cdot \mathsf{A}_1^{\text{QM}}(a_{pb}, \tilde{\eta}_{pb}) , \qquad (A22)$$

where the arguments of the function are defined by

$$a_{pb} = \frac{1}{2} \beta_b m_b v_p^2$$

$$\tilde{\eta}_{pb} = \frac{e_p e_b}{4\pi \hbar v_p} = |Z_p Z_b| \frac{e^2}{8\pi a_0} \frac{2a_0}{\hbar} \frac{1}{v_p} = |Z_p Z_b| \cdot 13.606 \,\text{eV} \cdot \frac{2 \cdot 5.29 \times 10^{-9} \,\text{cm}}{6.5821 \times 10^{-16} \,\text{eV} \,\text{s}} \frac{1}{v_p}$$

$$= 2.1870 \times 10^8 \frac{|Z_p Z_b|}{v_p \cdot (\text{cm/s})^{-1}} ,$$
(A24)

and the function itself takes the form

$$\mathsf{A}_{1}^{\text{QM}}(a,\eta) = -\int_{0}^{\infty} du \, \left[ \operatorname{Re} \psi \left\{ 1 + i \, \frac{\eta}{u} \right\} - \ln \left\{ \frac{\eta}{u} \right\} \right]$$

$$\frac{1}{2a \, u} \left[ \left( e^{-a(u-1)^{2}} + e^{-a(u+1)^{2}} \right) - \frac{e^{-a(u-1)^{2}} - e^{-a(u+1)^{2}}}{2a \, u} \right] \, . \tag{A25}$$

acoeff.f90:

END SUBROUTINE a\_quantum

```
am =-au-au2-a
ch =0.5*(EXP(ap)+EXP(am))
     sh = 0.5*(EXP(ap)-EXP(am))
  ENDIF
daq=-psilog*2*(ch - sh/au)/au
END FUNCTION daq
SUBROUTINE a_quantum(ib, a, eta, aq) IMPLICIT NONE
  INTEGER, INTENT(IN)
                                       ! species index
                           :: ib
                                       ! [dimensionless] (1/2) betab mb vp^2
  REAL,
             INTENT(IN)
                           :: a
  REAL,
                                       ! [dimensionless] ep eb/4pi hbar vp
             INTENT(IN)
                            :: eta
  REAL,
             INTENT(OUT) :: aq
                         :: u0, u1, du, u, um
  REAL
  INTEGER, PARAMETER :: NQ=1000 ! integration regions quantum : must REAL, PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad
  REAL,
  REAL,
             PARAMETER :: W13=0.555555556E0, W2=0.8888888889E0
            :: daq
  REAL
  INTEGER :: iu
  aq=0
u0=0.
  aq=0
  IF (ib == 1) THEN
      u0=0
      u1=4./SQRT(a)
  ELSE
      u0=1-10./SQRT(a)
      u0=MAX(0.,u0)
u1=1+10./SQRT(a)
  ENDIF
  du=(u1-u0)/NQ
  u=u0-du
  DO iu=1,NQ,2 ! Gaussian quadrature u=u+2.E0*du
      aq=aq+W2*daq(u,a,eta)
      um=u-du*UPM
      aq=aq+W13*daq(um,a,eta)
      um=u+du*UPM
      aq=aq+W13*daq(um,a,eta)
  ENDDO
  aq=aq*du
```

## Appendix B: Complete Source Code Listing: acoeff.f90

This section gives the complete listing for the source acoeff.f90 as it currently stands, including comments for the user and comments that I have made for myself (the latter will eventually disappear). This source module calculates the A-coefficients (coeff\_bps), their low energy asymptotic limits (coeff\_bps\_small\_E), and their high energy asymptotic limits (coeff\_bps\_high\_E). For the electron, there are two distinct high energy regions: an intermediate high energy regime  $T \ll E_p \ll (m_{\rm I}/m_e) T$  and an extreme high energy regime  $E \gg (m_{\rm I}/m_e) T$ .

Note: Currently, the subroutine coeff\_bps\_high\_E only returns the total electron contribution  $\mathcal{A}_e$  (regular + singular + quantum).

To do: To complete the subroutine coeff\_bps\_high\_E, I need to calculate the regular and singular contributions for the electrons in both high energy regimes. That is to say, I need to calculate the asymptotic limits of  $\mathcal{A}_{e,s}^{c}$  and  $\mathcal{A}_{e,s}^{c}$  respectively. The quantum asymptotic correction can then be obtained by

$$\mathcal{A}_e^{\mathrm{QM}} = \mathcal{A}_e - \mathcal{A}_e^{\mathrm{C}} = \mathcal{A}_e - (\mathcal{A}_{e-S}^{\mathrm{C}} + \mathcal{A}_{e-B}^{<}).$$

Before coeff\_bps\_high\_E returns the same quantities for electrons and ions, I will need to analytically calculate  $\mathcal{A}_{e,s}^{c}$  and  $\mathcal{A}_{e,R}^{c}$  and their high energy limits. To repeat the above note: until I finish this calculation, the subroutine coeff\_bps\_high\_E only returns the total electron contribution  $\mathcal{A}_{e}$ , although it returns the complete set of contributions for  $\mathcal{A}_{I}$ .

## 1. The A-Coefficients

acoeff.f90:

```
Robert Singleton
- Santa Fe, Winter 2005
- Santa Fe, March 2009 [start rewrite]
- Santa Fe, November 2009 [finish rewrite]

ROUTINE: bps_acoeff_ab_mass(nni, ep, mp, zp, ia, ib, betab, zb, mb, nb, & a_ab, a_ab_sing, a_ab_reg, a_ab_qm)

Assume a plasma composed of several species b, each separately in thermal equilibrium with themselves but not necessarily with each other[1]. This routine returns several useful components of the corresponding A-coefficients introduced in Note [2] below (BPS).

UNITS: A_{pb} has units of [MeV/micron] (subject to change in updates)
```

ı

[2] BPS paper

```
THE PHYSICS:
! The various subsystems b will exchange coulomb energy and they will eventually equilibrate to a common temperature. The A-coefficients
 introduced in Ref. [2] encode this coulomb energy exchange, exactly to
! leading and next-to-leading orders in the plasma coupling constant g.
! See Refs. [3,4,5] for more details. For a weakly coupled plasma (g << 1),
! the BPS calculation is essentially exact, and the error is O(g). Physical
 properties of interest, such as the stopping power dE/dx and the temperature
 equilibration rate between plasma species, can be obtained directly from
 the A-coefficients.
 Since electrons are thousands of times lighter than ions, one of the most
 physically accessible regime is the in which the electrons have a temperature
 T_e and the ions have a (possibly different) common temperature T_I. This
! is why the output is organized into electron contributions and total ion
 contributions (sum over all ions).
 INPUT: nni, ep, zp, mp, ia, ib, betab, zb, mb, nb
 Describe the incident projectile and the background plasma.
 projectile input quantities:
! ep : classical kinetic energy of the projectile [keV]
 zp : charge of the projectile in units of Z_p [dimensionless]
 mp : mass of the projectile [keV], i.e. mp = mp[grams]*c^2
 plasma input quantities:
      : Number of total plasma species = number ion species + 1
 nni
       : Charges of the plasma species. By convention zp(1) is the
 zb
         electron plasma component. [dimensionless, Array]
 betab: Inverse temperatures of the plasma components. For an
         electron-ion plasma, set betab(1)=1/T_e and all other
         values of the array to 1/T_I [keV^-1].
ı
 mb
       : Masses of the plasma species [keV].
       : Number densities of the plasma species [cm^-3].
       : First plasma species [usually the projectile]
 iа
       : Second plasma species
 OUTPUT: a_ab, a_ab_sing, a_ab_reg, a_ab_qm
! Each plasma component b makes a linear contribution A_b to the total
 A-coefficient, i.e. A = sum_b A_b [5]. Each A_b in turn can be be
 decomposed into a classical-quantum or electron-ion contributions.
                      : ac_e
: ac_i [sum over all ions]
 classical electron
 classical ion
 classical total
                       : ac_tot = ac_e + ac_i
! quantum
            electron
                       : aq_e
                       : aq_i [sum over all ions]
! quantum
                       : a_tot = aq_e + aq_i
 quantum
            total
                       : a_e = ac_e + aq_e
: a_i = ac_i + aq_i
 total
            electric
 total
            ion
                       ! a_tot = a_e + a_i
 total
 NOTES:
  [1] The temperatures T_b may therefore all differ. By convention
      I take b=1 for the electron component of the plasma. A very
      useful and interesting parameter regime is the one in which the ions have a common temperature T\_I and the electron have
      a temperature T_e, usually with T_e = /= T_I. See also USAGE
      and note [3] below.
```

- L. Brown, D. Preston, and R. Singleton Jr., "Charged Particle Motion in a Highly Ionized Plasma", Physics Reports, 410 (2005) 237 [arXiv:physics/0501084]
- [3] The code employs rationalized cgs units in which the dimensionless plasma coupling parameter is defined by  $g = e^2 \text{ kappa/(4Pi*T)}$ ; in these units the Debye wavenumber is determined by  $\text{kappa}^2 = e^2 \text{ n/T}$  and the plasma frequency by  $\text{omega}^2 = e^2 \text{ n/m}$ . A weakly coupled plasma is one for which g << 1, i.e. a plasma with thermal kinetic energy (of order the temperature T) dominates the coulomb potential energy (for particles separated by a Debye length). In the more common non-rationalized cgs units, we define  $g = e^2 \text{ kappa/T}$ , with  $\text{kappa}^2 = 4 \text{ Pi } e^2 \text{ n/T}$  and  $\text{omega}^2 = 4 \text{ Pi } e^2 \text{ n/m}$ .
- [4] For coulomb energy exchange processes, the leading and next-to-leading order terms in the plasma coupling g are proportional to  $g^2*ln(g)$  and  $g^2$ , respectively. That is to say, for a property denoted by F, one can expand F in powers of g in the form:

$$F(g,eta) = A(eta)*g^2*ln(g) + B(eta)*g^2 + O(g^3)$$
  
=  $A(eta)*g^2*[ln(C(eta)*g)+O(g)],$ 

where eta is the dimensionless quantum parameter (eta <<1 means extreme classical scattering while eta >>1 means the extreme quantum limit). The relative error of BPS is therefore O(g). At the center of the sun g=0.4, and so the error of Ref. [1] is only of order 4% in this case. For the processes of charged stopping power and electronion temperature equilibration, Ref. [1] calculates the corresponding functions A(eta) and B(eta) exactly, including all orders in the two-body quantum scattering parameter eta =  $e^2/4$ Pi\*hbar\*v\_thermal (this means that BPS gives the correct interpolation between the classical and quantum regimes, exact to leading and next-to-leading order). The  $O(g^3)$  terms physically correspond to 3-body correlations within the plasma, and for a sufficiently weak plasma these are negligible. For strongly coupled plasmas (g >> 1), all terms in a g-expansion are important and the BPS calculation is not applicable.

[5] It makes sense to talk about separate linear contribution A\_b contributing \*from\* a given plasma component b only for a weakly coupled plasma. More exactly, A=sum\_b A\_b holds true only up to leading and next-to-leading order in the plasma coupling g. This is the order to which Ref. [2] calculates all quantities, and therefore BPS works to a consistent order in g.

where e^2

```
- = 13.606E-3
                                 [keV]
 Ве
        8 Pi a0
                                 Bohr radius: a0=5.29E-9 cm
SUBROUTINE bps_acoeff_ab_mass(nni, ep, mp, zp, ia, ib, betab, zb, mb, nb, &
            a_ab, a_ab_sing, a_ab_reg, a_ab_qm)
     USE physvars
     USE mathvars
       IMPLICIT NONE
                                                                     Plasma:
                                             INTENT(IN)
       INTEGER,
                                                                      number of ions
                                                         :: nni
       REAL,
                                            INTENT(IN)
                                                         :: ep
                                                                      energy input [ke
       REAL,
                                            INTENT(IN)
                                                         :: mp
                                                                      mass [keV]
                                            INTENT(IN)
       REAL,
                                                         :: zp
                                                                      charge
                                             INTENT(IN)
       INTEGER,
                                                         :: ia
                                            INTENT(IN)
       INTEGER,
                                                            ib
       REAL,
                 DIMENSION(1:nni+1),
                                            INTENT(IN)
                                                         :: betab
                                                                      temp array [1/ke
       REAL,
                DIMENSION(1:nni+1),
                                                                      mass array [keV]
                                            INTENT(IN)
                                                         :: mb
       REAL,
                 DIMENSION(1:nni+1),
                                            INTENT(IN)
                                                                      density [1/cc]
                                                         :: nb
       REAL.
                DIMENSION(1:nni+1),
                                            INTENT(IN)
                                                         :: zb
                                                                      charge array
                                                                    A-coeffs [MeV/mic
       REAL,
                                            INTENT(OUT) :: a_ab
                                             INTENT(OUT) :: a_ab_sing
       REAL,
       REAL.
                                            INTENT(OUT) :: a_ab_reg
       REAL,
                                            INTENT(OUT) :: a_ab_qm
                                     :: mpb, mbpb, kb2, ab
:: vp, zp2, k, k2, kd, kd2, a, b, eta
       REAL,
                 DIMENSION(1:nni+1)
       REAL
       REAL
                                     :: ac_r, ac_s, aq, c1, c2
                                     :: EPS_SMALL_E=2.E-4
:: EPS_SMALL_E_SING=2.E-4
       REAL, PARAMETER REAL, PARAMETER
       REAL, PARAMETER
                                     :: EPS_SMALL_E_REG=2.E-4
 initialize components of A-coefficients
       kb2=8*PI*AOCM*BEKEV*zb*zb*nb*betab
                                        [1/cm<sup>2</sup>]
[1/cm]
[1/cm<sup>2</sup>]
[1/cm]
       kd2 = SUM(kb2)
       kd = SQRT(kd2)
           = kb2(1)= SQRT(k2)
       k2
                                                 k = k_e
 Loop over charged plasma species
       mpb = mp*mb/(mp+mb)
                                       ! [keV]
                                       ļ
                                         [dimensionless]
       mbpb= mb/mpb
       vp =CC*SQRT(2*ep/mp)
                                       ļ
                                         [cm/s]
       zp2=zp**2
                                         [dimensionless]
                                         ab=(1/2) betab(ib)*mbc2(ib)*vp2/CC2
       ab =0.5*betab*mb*vp*vp/CC2
                                       ! [dimensionless]
       IF (zb(ib) .NE. O.) THEN
          =ab(ib)
          =-Log(2*betab(ib)*BEKEV*ABS(zp*zb(ib))*k*AOCM*mbpb(ib) )-2*GAMMA+2
       eta=ABS(zp*zb(ib))*2.1870E8/vp ! defined with projectile velocity vp
       c1=2*zp2*BEKEV*kb2(ib)*AOCM
                                       ! [keV/cm] c1 = e_p^2 kappa_b^2/(4 Pi)
       c1=c1*1.E-7
                                         [MeV/micron]
       c2=SQRT(a/PI)
                                         [dimensionless]
                                       ! c2=SQRT(betab(ib)*mb(ib)/TWOPI)*vp/CC
! A_{ab}-classical-singular
```

```
ļ
        CALL a_sing_mass(a,b,ac_s)
        a_ab_sing=c1*c2*ac_s
 A_{ab}-classical-regular
        CALL a_reg_mass(nni,ia,ib,vp,k2,kb2,betab,mb,ac_r)
        a_ab_reg=c1*ac_r
 A_{ab}-quantum
        CALL a_quantum_mass(ia,ib,a,eta,aq) ! eta = dimensionless quantum param.
        a_ab_qm=c1*c2*aq
 A_{ab}-total
        a_ab=a_ab_sing + a_ab_reg + a_ab_qm
        ENDIF
      END SUBROUTINE bps_acoeff_ab_mass
SUBROUTINE bps_acoeff_ab_matrix(nni, ep, betab, zb, mb, nb,
        a_ab, a_ab_sing, a_ab_reg, a_ab_qm, a_tot, a_i, a_e, ac_tot, &
        ac_i, ac_e, aq_tot, aq_i, aq_e, ac_s_i, ac_s_e, ac_r_i, ac_r_e)
     USE physvars
USE mathvars
        IMPLICIT NONE
                                                                     Plasma:
        INTEGER,
                                                         :: nni
                                             INTENT(IN)
                                                                      number of ions
        REAL,
                                             INTENT(IN)
                                                         :: ep
                                                                      energy
        REAL.
                 DIMENSION(1:nni+1),
                                             INTENT(IN)
                                                         :: betab
                                                                      temp array [1/ke
        REAL.
                 DIMENSION(1:nni+1),
                                            INTENT(IN)
                                                         :: zb
                                                                      charge array
        REAL,
                 DIMENSION(1:nni+1),
                                            INTENT(IN)
                                                         :: mb
                                                                      mass array [keV]
        REAL,
                 DIMENSION(1:nni+1),
                                            INTENT(IN)
                                                         ::
                                                           nb
                                                                      density [1/cc]
                                                                    A-coeffs [MeV/mic
                 DIMENSION(1:nni+1,1:nni+1),INTENT(OUT) :: a_ab
       REAL,
                 DIMENSION(1:nni+1,1:nni+1),INTENT(OUT) :: a_ab_sing
        REAL.
        REAL.
                 DIMENSION(1:nni+1,1:nni+1),INTENT(OUT) :: a_ab_reg
        REAL,
                 DIMENSION(1:nni+1,1:nni+1),INTENT(OUT) :: a_ab_qm
        REAL,
                 DIMENSION(1:nni+1),
                                                         :: a_tot
                                             INTENT (OUT)
                                            INTENT(OUT)
INTENT(OUT)
INTENT(OUT)
                 DIMENSION(1:nni+1),
DIMENSION(1:nni+1),
DIMENSION(1:nni+1),
        REAL,
                                                         :: a_i
        REAL,
                                                         :: a_e
        REAL,
                                                         ::
                                                           ac_tot
                 DIMENSION(1:nni+1),
        REAL,
                                             INTENT (OUT)
                                                        :: ac_i
       REAL,
                 DIMENSION(1:nni+1),
                                             INTENT (OUT)
                                                        :: ac_e
        REAL,
                 DIMENSION(1:nni+1),
                                             INTENT (OUT)
                                                        :: aq_tot
        REAL,
                 DIMENSION(1:nni+1),
                                             INTENT(OUT) :: aq_i
                 DIMENSION(1:nni+1),
        REAL,
                                             INTENT(OUT) :: aq_e
        REAL,
                 DIMENSION(1:nni+1),
                                             INTENT (OUT)
                                                        :: ac_s_i
                                            INTENT(OUT)
INTENT(OUT)
        REAL,
                 DIMENSION(1:nni+1),
                                                        :: ac_s_e
        REAL,
                 DIMENSION(1:nni+1)
                                                         :: ac_r_i
                 DIMENSION(1:nni+1),
                                            INTENT(OUT) :: ac_r_e
        REAL,
        REAL
                :: aab, aab_sing, aab_reg, aab_qm
        REAL
        REAL :: mp, zp
INTEGER :: ia, ib
        a_i
        ac_s_i = 0
        ac_r_i = 0
             = 0
        ac_i
        aq_i
             = 0
        DO ia=1,nni+1
```

```
mp=mb(ia)
          zp=zb(ia)
          \overline{D}0 ib=1,nni+1
            CALL bps_acoeff_ab_mass(nni, ep, mp, zp, ia, ib, betab, zb, mb, nb, &
            aab, aab_sing, aab_reg, aab_qm)
            a_ab(ia,ib)
                             =aab
            a_ab_sing(ia,ib)=aab_sing
            a_ab_reg(ia,ib) =aab_reg
            a_ab_qm(ia,ib)
                             =aab_qm
            IF (ib == 1) THEN
               a_e(ia)
                          = aab
               ac_s_e(ia) = aab_sing
                ac_r_e(ia)= aab_reg
               ac_e(ia) = aab_sing + aab_reg
               aq_e(ia)
                         = aab_qm
            ELSE
               a_i(ia)
                          = a_i(ia)
                                        + aab
               ac_s_i(ia) = ac_s_i(ia) + aab_sing
               ac_r_i(ia) = ac_r_i(ia) + aab_reg
               ac_i(ia) = ac_i(ia)
                                        + aab_sing + aab_reg
                aq_i(ia) = aq_i(ia)
                                        + aab_qm
            ENDIF
          ENDDO
        a_tot(ia) = a_e(ia) + a_i(ia)
ac_tot(ia)= ac_e(ia) + ac_i(ia)
aq_tot(ia)= aq_e(ia) + aq_i(ia)
ENDDO
      END SUBROUTINE bps_acoeff_ab_matrix
! Returns A_{p I} = \sum_i A_{p i} for backward compatibility
SUBROUTINE bps_acoeff_ei_mass(nni, ep, zp, mp, betab, zb, mb, nb, & a_tot, a_i, a_e, ac_tot, ac_i, ac_e, aq_tot, aq_i, aq_e,&
            ac_s_i, ac_s_e, ac_r_i, ac_r_e)
     USE physvars
USE mathvars
      USE controlvars
        IMPLICIT NONE
                                                               ! Plasma:
        INTEGER,
                                       INTENT(IN)
                                                    :: nni
                                                                  number of ions
                 DIMENSION(1:nni+1), INTENT(IN)
                                                    :: betab
                                                                  temp array [1/keV]
        REAL,
                 DIMENSION(1:nni+1), INTENT(IN)
                                                                  mass array [keV]
        REAL,
                                                    :: mb
        REAL,
                 DIMENSION(1:nni+1), INTENT(IN)
                                                    :: nb
                                                                  density [1/cc]
        REAL,
                 DIMENSION(1:nni+1), INTENT(IN)
                                                    :: zb
                                                                  charge array
                                                                 Projectile
        REAL,
                                       INTENT(IN)
                                                                  projectile energy [keV]
                                                    :: ep
        REAL,
                                       INTENT(IN)
                                                                  projectile mass
                                                                                      [keV]
                                                    :: mp
        REAL,
                                       INTENT(IN)
                                                    :: zp
                                                                  projectile charge
                                                                 A-coeffs [MeV/micron]
        REAL,
                                       INTENT(OUT)
                                                                  electron + ion
                                                   :: a_tot
                                       INTENT (OUT)
INTENT (OUT)
INTENT (OUT)
INTENT (OUT)
        REAL,
                                                    :: a_i
                                                                  ion contribution
        REAL,
                                                    :: a_e
                                                                  electron contribution
        REAL,
                                                    :: ac_tot
                                                                  classical
        REAL,
                                                    :: ac_i
                                                                  classical
        REAL,
                                       INTENT(OUT)
                                                   :: ac_e
                                                                  classical
                                       INTENT(OUT) :: aq_tot
        REAL,
                                                                  quantum
                                       INTENT(OUT) :: aq_i
        REAL.
                                                                  quantum
        REAL,
                                       INTENT(OUT) :: aq_e
                                                                  quantum
        REAL,
                                       INTENT(OUT) :: ac_s_i
                                       INTENT(OUT) :: ac_s_e
INTENT(OUT) :: ac_r_i
        REAL,
        REAL,
```

```
REAL,
                                         INTENT(OUT) :: ac_r_e
                 :: adum, ac_s, ac_r, aq
:: ia, ib, nnb
        REAL
        INTEGER
  initialize components of A-coefficients
        a_{tot} = 0
                    ! electron + ion
               =0
                   !
                      ion contribution
        a_i
        a_e
               =0
                      electron contribution
        ac_tot=0
                     classical total
                    ! classical electron
        ac_e = 0
        ac_i
              =0
                    ! classical ion
        aq_tot=0
                    ! quantum total
        aq_e =0
aq_i =0
                    ! quantum electron
                    ! quantum ion
        a\bar{c}_s=0
        ac_s_e=0
ac_r_i=0
ac_r_e=0
        NNB = nni+1
                                        ! number of ions + electrons
        ia=1
        DO ib=1,nni+1
           (zb(ib) .NE. O.) THEN
             CALL bps_acoeff_ab_mass(nni, ep, mp, zp, ia, ib, betab, zb, mb, nb, &
             adum, ac_s, ac_r, aq)
            CALL x_collect(ib, NNB, ac_s, ac_r, aq, a_tot, a_i, a_e, ac_tot, ac_i, ac_e, aq_tot,
             aq_i, aq_e, ac_s_i, ac_s_e, ac_r_i, ac_r_e)
        ENDIF
        ENDDO
      END SUBROUTINE bps_acoeff_ei_mass
! singular contribution for non-zero electron mass
SUBROUTINE a_sing_mass(a, b, ac_s)
                  INTENT(IN)
INTENT(IN)
        REAL,
                                :: a
        REAL,
                                :: b
                               :: ac_s
        REAL,
                  INTENT(OUT)
        REAL :: u0, u1, du, u, um

INTEGER, PARAMETER :: NS=1000 ! integration regions: must be even

REAL, PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad

REAL, PARAMETER :: W13=0.555555556E0, W2=0.8888888889E0
           ac_s=0
           u0=0
           u1 = 1
           du=(u1-u0)/NS
           u=u0-du
           DO iu=1,NS,2 ! Gaussian quadrature
               u=u+2.E0*du
               ac_s=ac_s+W2*dab_sing(u,a,b)
               um=u-du*UPM
               ac_s=ac_s+W13*dab_sing(um,a,b)
               um=u+du*UPM
               ac_s=ac_s+W13*dab_sing(um,a,b)
           ENDDO
            ac_s=ac_s*du
      END SUBROUTINE a_sing_mass
!
      FUNCTION dab_sing(u, a, b)
        IMPLICIT NONE
        REAL,
                       INTENT(IN)
                                                    [dimensionless]
                                                    [dimensionless]
        REAL,
                       INTENT(IN)
                                    :: a
                                                  ! a=(1/2)*beta*mpc2*vp^2/C^2
```

```
:: b ! [dimensionless]
:: dab_sing ! [dimensionless]
       REAL,
                    INTENT(IN)
       REAL
       dab\_sing=SQRT(u)*EXP(-a*u)*(-LOG(u/(1-u)) + b)
     END FUNCTION dab_sing
! regular contribution for non-zero electron mass
FUNCTION dab_reg(u, vp, ia, ib, nni, k2, kb2, betab, mb)
     USE mathvars
     USE physvars IMPLICIT NONE
       REAL,
                                    INTENT(IN)
                                                             [dimensionless]
                                                 :: u
                                                :: vp
                                    INTENT(IN)
       REAL,
                                                           ! Projectile velocity [cm/
       INTEGER,
                                    INTENT(IN)
                                                :: ia
                                                           !
                                                            Species number
       INTEGER,
                                    INTENT(IN)
                                                           ! Species number
                                                :: ib
       INTEGER,
                                    INTENT(IN)
                                                :: nni
                                                           ! Number of ion species
       REAL,
                                    INTENT(IN)
                                                :: k2
                                                           ! Wave-number squared [1/c
       REAL,
                DIMENSION(1:nni+1), INTENT(IN)
                                                :: kb2
                                                           ! Debye wavenumber squared
                DIMENSION(1:nni+1), INTENT(IN)
       REAL,
                                                          ! Temperature array [1/keV
                                                :: betab
       REAL,
                DIMENSION(1:nni+1), INTENT(IN)
                                                           ! Mass array [keV]
                                                :: mb
       R.E.A.L.
                                                 :: dab_reg! [dimensionless]
       REAL,
                DIMENSION(1:nni+1) :: kbar2b, ab, ab2
       REAL
                                    :: fr, fi, fabs, farg, h, r_ib
       REAL
                                    :: kcb, bm_ic, bm_ib, a_ic, a_ib, ex, au
       INTEGER
                                    :: ic
       ab=SQRT(0.5*betab*mb)*vp/CC
       ab2=ab*ab
       kbar2b=kb2/k2
       CALL frfi(u,nni,kbar2b,ab,fr,fi,fabs,farg)
       h=2*(fr*farg + fi*LOG(fabs))*u
!*!
        h=2*(fr*farg + fi*LOG(fabs))
1 * 1
 construct spectral weight ratio Rb=rho_b/rho_tot
!*!
         r_ib=0
         DO ic=1,nni+1
           r_ib=r_ib + kbar2b(ib)*(ab(ic)/ab(ib))*EXP((ab2(ib)-ab2(ic))*u*u)
         ENDDO
         r_{ib}=1./r_{ib}
!*!
       r_{ib}=0
       bm_ib=betab(ib)*mb(ib)
       a_{ib} = ab(ib)*ab(ib)
       D0 ic=1,nni+1
          kcb=kb2(ic)/k2
          bm_ic=betab(ic)*mb(ic)
          a_{ic} = ab(ic)*ab(ic)
          \overline{IF} (ic == ib) THEN
             ex=1.
          ELSE
             au=(a_ic-a_ib)*u
             ex=EXP(-au)
          ENDIF
          r_ib=r_ib + kcb*SQRT(bm_ic/bm_ib)*ex
       ENDDO
       r_{ib}=1./r_{ib}
!*!
!r_{ib}=1.
!*!
ļ
       dab_reg=-r_ib*h/TWOPI
```

```
END FUNCTION dab_reg
      SUBROUTINE a_reg_mass(nni, ia, ib, vp, k2, kb2, betab, mb, ac_r)
      USE physvars
IMPLICIT NONE
                                          INTENT(IN)
INTENT(IN)
         INTEGER,
                                                         :: nni
         INTEGER,
                                                        :: ia
         INTEGER,
                                           INTENT(IN)
                                                        :: ib
                                          INTENT(IN)
        REAL,
                                                        :: vp
        REAL,
                                                         :: k2
                                          INTENT(IN)
                                          INTENT(IN)
INTENT(IN)
INTENT(IN)
INTENT(OUT)
         REAL,
                                                        :: kb2
                   DIMENSION(1:nni+1),
         REAL,
                                                        :: betab
         REAL,
                   DIMENSION(1:nni+1),
                                                        :: mb
         REAL,
                                                        :: ac_r
        REAL,
INTEGER,
                   DIMENSION(1:nni+1)
                                           :: ab
                   PARAMETER :: NR=10 ! integration regions: must be even
PARAMETER :: NR=100 ! integration regions: must be even
PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad
PARAMETER :: W13=0.5555555556E0, W2=0.8888888889E0
:: u0, u1, du, u, um, dab_reg
ļ
         INTEGER,
         REAL,
        REAL,
         R.E.A.L.
         INTEGER
                                  iu
         ab=SQRT(0.5*betab*mb)*vp/CC
         ac_r=0
         u0=0.0
         u1=1
ļ
         u1=MIN(1.,5/(ab(ib)**2))! support can lie << 1
         du=(u1-u0)/NR
         u=u0-du
         DO iu=1,NR,2 ! Gaussian quadrature
            u=u+2.*du
            ac_r=ac_r+W2*dab_reg(u,vp,ia,ib,nni,k2,kb2,betab,mb)
            um=u-du*UPM
            ac_r=ac_r+W13*dab_reg(um,vp,ia,ib,nni,k2,kb2,betab,mb)
            um=u+du*UPM
            ac_r=ac_r+W13*dab_reg(um,vp,ia,ib,nni,k2,kb2,betab,mb)
         ENDDO
         ac_r=ac_r*du
!*!
          11m = 1
          ac_r=dab_reg(um, vp, ia, ib, nni, k2, kb2, betab, mb)
!*!
      END SUBROUTINE a_reg_mass
! quantum contribution for non-zero electron mass
FUNCTION daq(u, a, eta)
      USE physvars IMPLICIT NONE
                                          INTENT(IN)
INTENT(IN)
         REAL,
                                                                            [dimensionless]
                                                        :: u
        REAL,
                                                        :: a
                                                                            [dimensionless]
         REAL,
                                          INTENT(IN)
                                                        :: eta
                                                                           [dimensionless]
                                                                  ! [dimensionless]
         REAL
                                                        :: daq
         REAL
                           :: repsi, au, eu, au2, ap, am, psilog, ch, sh, csh
         eu=eta/u
         psilog=repsi(eu) - LOG(eu)
         au =2*a*u
         au2=a*u*u
         ap = au - au2 - a
         am =-au-au2-a
ch =0.5*(EXP(ap)+EXP(am))
         sh = 0.5*(EXP(ap)-EXP(am))
         csh=2*(ch - sh/au)/au
         daq=-psilog*csh
      END FUNCTION dag
```

```
SUBROUTINE a_quantum_mass(ia, ib, a, eta, aq) IMPLICIT NONE
                         INTEGER, INTENT(IN)
                                                                                             :: ia
                                                                                                                           ! species index
                         INTEGER, INTENT(IN)
                                                                                           :: ib
                                                                                                                          ! species index
                                                                                                                         ! [dimensionless] (1/2) betab mb vp^2
                                                                                          :: a
                         REAL,
                                                      INTENT(IN)
                         REAL,
                                                      INTENT(IN)
                                                                                            :: eta
                                                                                                                     ! [dimensionless] ep eb/4pi hbar vp
                         REAL,
                                                      INTENT(OUT) :: aq
                                                                                       :: u0, u1, du, u, um
                         REAL
                                                      PARAMETER :: NQ=1000 ! integration regions quantum : must PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad
                         INTEGER, PARAMETER :: NQ=1000
                         REAL,
                         REAL,
                                                     PARAMETER :: W13=0.555555556E0, W2=0.8888888889E0
                         REAL
                                                   :: daq
                         INTEGER :: iu
                         aq=0
                         u0=0.
                         aq=0
                         IF (ib == ia) THEN
                                  u0 = 0
                                  u1=4./SQRT(a)
                         ELSE
                                  u0=1-10./SQRT(a)
                                  u0=MAX(0.,u0)
u1=1+10./SQRT(a)
                         ENDIF
                         du=(u1-u0)/NQ
                         u=u0-du
                         DO iu=1,NQ,2 ! Gaussian quadrature
u=u+2.E0*du
                                  aq=aq+W2*daq(u,a,eta)
                                  um=u-du*UPM
                                   aq=aq+W13*daq(um,a,eta)
                                  um=u+du*UPM
                                   aq=aq+W13*daq(um,a,eta)
                         ENDDO
                 aq=aq*du
END SUBROUTINE a_quantum_mass
!====== dE/dx from A-coefficient =======
    **XX** !
     This is a driver to check the analytic evalulation dE_b/dx
     against the one obtained by differentiating A_b.
                                            [ 1 - ----- Sum_l ----- {\hat vp}^l A_b(vp) ] 
 [ beta_b mp vp d vp^l ]
                                  = \begin{bmatrix} 1 & - & \frac{2}{1} & \frac{T_b}{1} & \frac{1}{1} & \frac{2}{1} & \frac{T_b}{1} & \frac{dA_b}{1} & \frac
                                   = \begin{bmatrix} 1 & - & T_b \\ 1 & - & --- \end{bmatrix} * A_b(Ep) - T_b * \frac{dA_b}{---}(Ep) 
                                     JTINE acoeff_dedx_bps(nni,ep,zp,mp,betab,zb,mb,nb, &
  dedx_a_tot, dedx_a_i, dedx_a_e, dedxc_a_tot, dedxc_a_i, dedxc_a_e, &
  dedxq_a_tot, dedxq_a_i, dedxq_a_e, dedxc_a_s_i, dedxc_a_s_e, &
                  SUBROUTINE acoeff_dedx_bps(nni,ep,zp,mp,betab,zb,mb,nb,
                                     dedxc_a_r_i, dedxc_a_r_e)
                  USE physvars
USE mathvars
                         IMPLICIT NONE
                                                                                                                                                                                                ! Plasma:
                                                                                                                       INTENT(IN) :: nni
                         INTEGER,
                                                                                                                                                                                                    number of ions
```

```
REAL,
          DIMENSION(1:nni+1), INTENT(IN)
                                                :: betab
                                                               temp array [1/keV]
          DIMENSION(1:nni+1), INTENT(IN)
REAL,
                                                :: mb
                                                            ļ
                                                               mass array [keV]
          DIMENSION(1:nni+1), INTENT(IN)
REAL,
                                                            ļ
                                                               density [1/cc]
                                                :: nb
REAL,
          DIMENSION(1:nni+1), INTENT(IN)
                                                :: zb
                                                               charge array
                                                              Projectile
REAL,
                                  INTENT(IN)
                                                               projectile energy [keV]
                                                :: ep
REAL,
                                  INTENT(IN)
                                                               projectile mass
                                                                                     [keV]
                                                :: mp
REAL,
                                  INTENT(IN)
                                                               projectile charge
                                                :: zp
                                                                    dE/dx [MeV/micron]
REAL,
                                  INTENT (OUT)
                                                                     electron + ion
                                                :: dedx_a_tot
REAL,
                                  INTENT(OUT)
                                                :: dedx_a_i
                                                                      ion contribution
                                  INTENT(OUT)
INTENT(OUT)
INTENT(OUT)
REAL,
                                                                     electron contribut
                                                :: dedx_a_e
REAL,
                                                :: dedxc_a_tot
:: dedxc_a_i
                                                                     classical
REAL,
                                                                     classical
REAL,
                                  INTENT(OUT)
                                                :: dedxc_a_e
                                                                     classical
                                  INTENT(OUT) :: dedxq_a_tot
REAL,
                                                                     quantum
                                  INTENT(OUT) :: dedxq_a_i
REAL,
                                                                     quantum
REAL,
                                  INTENT(OUT) :: dedxq_a_e
                                                                     quantum
REAL,
                                                :: dedxc_a_s_i
                                  INTENT (OUT)
REAL,
                                  INTENT (OUT)
INTENT (OUT)
                                                :: dedxc_a_s_e
REAL,
                                                ::
                                                   dedxc_a_r_i
REAL.
                                  INTENT(OUT) :: dedxc_a_r_e
REAL :: a_tot_p, a_i_p, a_e_p, ac_tot_p, ac_i_p, ac_e_p, aq_tot_p, aq_i_p, aq
REAL :: ac_s_i_p, ac_s_e_p, ac_r_i_p, ac_r_e_p 

REAL :: a_tot_m, a_i_m, a_e_m, ac_tot_m, ac_i_m, ac_e_m, aq_tot_m, aq_i_m, aq
REAL :: ac_s_i_m, ac_s_e_m, ac_r_i_m, ac_r_e_m REAL :: a_tot, a_i, a_e, ac_tot, ac_i,
REAL :: a_tot, a_i, a_e, ac_to

REAL :: ac_s_i,ac_s_e, ac_r_i,ac_r_e

REAL :: te, ti, dep, dep2, epp, epm
                                                          ac_e,
                                                                   aq_tot,
                                                                               aq_i,
te =1./betab(1)
ti = 1./betab(2)
             = 0 ! electron + ion
dedx_a_tot
              = 0 !
dedx_a_i
                     ion contribution
              = 0 ! electron contribution
dedx_a_e
dedxc_a_tot = 0 ! classical
dedxc_a_i
              = 0 ! classical
dedxc_a_e = 0 !

dedxq_a_tot = 0 !
                  ! classical
                     quantum
dedxq_a_i
              = 0
                     quantum
              = 0
                  ! quantum
dedxq_a_e
dedxc_a_s_i =
dedxc_a_s_e =
                0
                0
dedxc_a_r_i = 0
dedxc_a_r_e = 0
dep = ep * 1.E-4

dep2 = 2 * dep
CALL bps_acoeff_ei_mass(nni,ep,zp,mp,betab,zb,mb,nb,a_tot,a_i, &
  a_e,ac_tot,ac_i,ac_e,aq_tot,aq_i,aq_e,ac_s_i,ac_s_e,&
  ac_r_i,ac_r_e)
epp = ep + dep
CALL bps_acoeff_ei_mass(nni,epp,zp,mp,betab,zb,mb,nb,a_tot_p,a_i_p, &
  a_e_p, ac_tot_p, ac_i_p, ac_e_p, aq_tot_p, aq_i_p, aq_e_p, ac_s_i_p, ac_s_e_p, &
  ac_r_i_p,ac_r_e_p)
epm = ep - dep
CALL bps_acoeff_ei_mass(nni,epm,zp,mp,betab,zb,mb,nb,a_tot_m,a_i_m, &
  a_e_m,ac_tot_m,ac_i_m,ac_e_m,aq_tot_m,aq_i_m,aq_e_m,ac_s_i_m,ac_s_e_m,&
  ac_r_i_m,ac_r_e_m)
```

aq

!

```
dedx_a_tot = (1 - te/ep)*a_tot - te*(a_tot_p - a_tot_m)/dep2
  dedx_a_i = (1 - ti/ep)*a_i - ti*(a_i_p - a_i_m)/dep2
  dedx_a_e = (1 - te/ep)*a_e - te*(a_e_p - a_e_m)/dep2
  dedxq_a_tot = (1 - te/ep)*aq_tot - te*(aq_tot_p-aq_tot_m)/(2*dep)
  dedxq_a_i = (1 - ti/ep)*aq_i - ti*(aq_i_p - aq_i_m)/dep2
                                 - te*(aq_e_p - aq_e_m)/dep2
  dedxq_a_e = (1 - te/ep)*aq_e
  dedxc_a_tot = (1 - te/ep)*ac_tot - te*(ac_tot_p-ac_tot_m)/(2*dep)
  dedxc_a_s_i = (1 - te/ep)*ac_s_i
                                      - te*(ac_s_i_p -ac_s_i_m)/dep2
  dedxc_a_s_e = (1 - te/ep)*ac_s_e
                                      - te*(ac_s_e_p - ac_s_e_m)/dep2
  dedxc_a_r_i = (1 - te/ep)*ac_r_i
                                      - te*(ac_r_i_p -ac_r_i_m)/dep2
  dedxc_a_r_e = (1 - te/ep)*ac_r_e
                                      - te*(ac_r_e_p -ac_r_e_m)/dep2
END SUBROUTINE acoeff_dedx_bps
SUBROUTINE a_collect(ib, ibmax, ac_s, ac_r, aq, a_tot, a_i, a_e, &
  ac_tot, ac_i, ac_e, aq_tot, aq_i, aq_e, ac_s_i, ac_s_e, ac_r_i, ac_r_e)
IMPLICIT NONE
  INTEGER, INTENT(IN)
                                    ! species index
                          :: ib
  INTEGER, INTENT(IN)
                          :: ibmax
                                   ! species index maximum = NNB+1
  REAL,
           INTENT(IN)
                          :: ac_s
                                    ! singular contribution
  REAL,
                         :: ac_r
           INTENT(IN)
                                    ! regular contribution
  REAL,
           INTENT(IN)
                          :: aq
                                    ! quantum contribution
 REAL,
           INTENT(INOUT) :: a_tot ! running total over ions
 REAL,
           INTENT(INOUT) :: a_i
                                   ! running total over ions
  REAL,
           INTENT(INOUT) :: a_e ! electron component
 REAL,
           INTENT(INOUT) :: ac_tot ! running total over ions
 REAL,
           INTENT(INOUT) :: ac_i
                                    ! running total over ions
 REAL,
           INTENT(INOUT) :: ac_e
                                    ! electron component
  REAL.
           INTENT(INOUT) :: aq_tot ! running total over ions
 REAL,
           INTENT(INOUT) :: aq_i
                                  ! running total over ions
 REAL,
           INTENT(INOUT) :: aq_e
                                    ! electron component
 REAL,
           INTENT(INOUT) :: ac_s_i
          INTENT(INOUT) :: ac_s_e
INTENT(INOUT) :: ac_r_i
INTENT(INOUT) :: ac_r_e
 REAL,
  REAL,
  REAL,
  REAL :: ac_sr
  ac_sr=ac_s + ac_r
IF (ib==1) THEN
     ac_e=ac_sr
     aq_e=aq
     a_e = ac_e + aq_e
     ac_s_e=ac_s
     ac_r_e=ac_r
  ELSE
     ac_i=ac_i + ac_sr
     aq_i=aq_i + aq
     a_{\overline{i}} = a_{\overline{i}} + ac_{\overline{s}} + aq
     ac_s_i=ac_s_i + ac_s
     ac_r_i=ac_r_i + ac_r
  ENDIF
  IF (ib==ibmax) THEN
     ac\_tot = ac\_e + ac\_i
     aq_tot = aq_e + aq_i
     a_{tot} = ac_{tot} + aq_{tot}
```

ENDIF END SUBROUTINE a\_collect

## 2. Low Energy Asymptotics

acoeff.f90 cont.:

```
ROUTINE: SUBROUTINE coeff_bps_small_E(nni, ep, zp, mp, betab, zb, mb, nb, & a_tot_lim, a_i_lim, a_e_lim, ac_tot_lim, ac_i_lim, ac_e_lim, aq_tot_lim,&
    aq_i_lim, aq_e_lim)
  The asymptotic low energy regime E_p << T: this routine returns several
  useful components of the corresponding A-coefficients in the low energy
  regime.
  UNITS: A_b has units of [MeV/micron] (subject to change in updates)
  The incident projectile and the background plasma.
! projectile input quantities:
! ep : classical kinetic energy of the projectile [keV]
  zp : charge of the projectile in units of Z_p [dimensionless]
  mp : mass of the projectile [keV], i.e. mp = mp[grams]*c^2
  plasma input quantities:
nni : Number of total plasma species = number ion species + 1
        : Charges of the plasma species. By convention zp(1) is the
  : electron plasma component. [dimensionless, Array] betab: Inverse temperatures of the plasma components. For an
           electron-ion plasma, set betab(1)=1/T_e and all other
           values of the array to 1/T_I.
ļ
         : Masses of the plasma species [keV].
  mb
         : Number densities of the plasma species [cm^-3].
  nb
  OUTPUT: a_tot_lim, a_i_lim, a_e_lim, ac_tot_lim, ac_i+lim, ac_e_lim,
            aq_tot_lim, aq_i_lim, aq_e_lim
! classical electron : ac_e_lim
                           : ac_i_lim [sum over all ions]
: ac_tot_lim = ac_e_lim + ac_i_lim
  classical ion classical total
! quantum
               electron
                           : aq_e_lim
! quantum
                            : aq_i_lim [sum over all ions]
               ion
               total
                            : a_tot_lim = aq_e_lim + aq_i_lim
! quantum
                            : a_e_lim = ac_e_lim + aq_e_lim
: a_i_lim = ac_i_lim + aq_i_lim
! a_tot_lim = a_e_lim + a_i_lim
               electric
! total
ļ
  total
ı
  total
       SUBROUTINE coeff_bps_small_E(nni, ep, zp, mp, betab, zb, mb, nb, a_tot_lim, a_i_lim, a_e_lim, ac_tot_lim, ac_i_lim, ac_e_lim, aq_tot_lim, aq_i_lim, aq_e_lim, ac_s_i_lim, ac_s_e_lim,
               ac_r_i_lim, ac_r_e_lim)
       USE physvars
USE mathvars
          IMPLICIT NONE
                     INTENT(IN) :: nn:
DIMENSION(1:nni+1), INTENT(IN) :: be-
DIMENSION(1:nni+1), INTENT(IN) :: mb
          INTEGER,
                                                               :: nni
                                                                                    number of ions
                                                               :: betab
          REAL,
                                                                                !
                                                                                    temp array [1/keV]
          REAL,
                                                                                    mass array [keV]
                     DIMENSION(1:nni+1), INTENT(IN) :: nb
          REAL,
                                                                                    density [1/cc]
```

```
REAL,
                   DIMENSION(1:nni+1), INTENT(IN)
                                                        :: zb
                                                                           charge array
         REAL,
                                          INTENT(IN)
                                                        :: ep
                                                                           projectile energy [k
         REAL,
                                          INTENT(IN)
                                                        :: mp
                                                                           projectile mass
         REAL,
                                          INTENT(IN)
                                                         :: zp
                                                                           projectile charge
                                                                          A-coeffs [MeV/micron]
                                          INTENT(OUT)
INTENT(OUT)
INTENT(OUT)
INTENT(OUT)
         REAL,
                                                        :: a_tot_lim
                                                                           electron + ion
         REAL,
                                                                           ion contribution
                                                        :: a_i_lim
         REAL,
                                                        :: a_e_lim
                                                                           electron contributio
         REAL,
                                                        :: ac_tot_lim!
                                                                           classical
         REAL,
                                           INTENT(OUT)
                                                        :: ac_i_lim
                                                                           classical
                                          INTENT(OUT)
INTENT(OUT)
         REAL,
                                                        :: ac_e_lim
                                                                           classical
         REAL,
                                                        :: aq_tot_lim!
                                                                           quantum
         REAL.
                                          INTENT (OUT)
                                                        :: aq_i_lim
                                                                           quantum
                                          INTENT(OUT)
         REAL,
                                                        :: aq_e_lim
                                                                           quantum
         REAL,
                                          INTENT (OUT)
                                                        :: ac_s_i_lim!
                                                                           singular
         REAL,
                                          INTENT(OUT) :: ac_s_e_lim!
                                                                           singular
         REAL,
                                          INTENT(OUT) :: ac_r_i_lim!
                                                                           regular
         REAL,
                                          INTENT(OUT) :: ac_r_e_lim!
                                                                           regular
         REAL
                  :: ac_r_lim, ac_s_lim, aq_lim
        INTEGER :: ib, nnb
REAL, DIMENSION(1:nni+1)
                                           :: kb2
                                                       \lfloor 1/\text{cm}^2 \rfloor
         REAL,
                   DIMENSION(1:nni+1)
                                          :: ab
                                                       [dimensionless]
         REAL
                  :: vp
                                                       [cm/s]
         REAL
                                                       [cm^2/s^2]
                  :: vp2
                                                       [1/cm^2]
         REAL
                  :: kd2
         REAL
                                                       [dimensionless]
                  :: a, zp2
         REAL
                                                       [keV/cm]
                  :: c1
         REAL
                  ::
                     c2
                                                       [dimensionless]
         nnb =nni+1
         vp = CC*SQRT(2*ep/mp)
                                            [cm/s]
                                            [cm^2/s^2]
         vp2 = vp*vp
         kb2 = DEBYE2*zb*zb*nb*betab
                                            [1/cm^2]
                                            [1/cm<sup>2</sup>]
         kd2 = SUM(kb2)
         zp2 = zp**2
                                            [dimensionless]
         ab =0.5*betab*mb*vp2/CC2
                                         ! [dimensionless]
                                         ! ab=(1/2) betab(ib)*mbc2(ib)*vp2/CC2
  initialize A-coefficients
        a_tot_lim =0
a_i_lim =0
                           electron + ion
                    =0
                           ion contribution
         a_e_lim
                    =0
                           electron contribution
         ac\_tot\_lim=0
                           classical total
         ac_e_lim ac_i_lim
                           classical electron classical ion
                   =0
                   =0
         aq_tot_lim=0
                           quantum total
                    =0
                           quantum electron
         aq_e_lim
                    =0
         aq_i_lim
                         ! quantum ion
         ac_s_i_lim=0
ac_s_e_lim=0
         ac_r_i_lim=0
         ac_r_e_lim=0
ļ
         DO ib=1,nni+1
         IF (zb(ib) .NE. O.) THEN
                                               [dimensionless] [keV/cm]
            a=ab(ib)
            c1=2*zp2*BEKEV*kb2(ib)*AOCM
                                            !
            c1=c1*1.E-7
                                               [MeV/micron]
            c2=SQRT(a/PI)
                                               [dimensionless]
  singular: asymptotic low energy form
```

[k

```
CALL a_sing_ib_small_E(nni,ib,ep,zp,mp,betab,zb,mb,nb,ac_s_lim)
ac_s_lim=c1*c2*ac_s_lim
 regular: asymptotic low energy form
           CALL a_reg_ib_small_E(nni,ib,ep,zp,mp,betab,zb,mb,nb,ac_r_lim)
           ac_r_lim=c1*ac_r_lim
ı
  quantum: asymptotic low energy form
          CALL aq_ib_small_E(nni, ib, ep, zp, mp, betab, zb, mb, nb, aq_lim)
          aq_lim=c1*c2*aq_lim
  collect components
           ac_s_i_lim, ac_s_e_lim, ac_r_i_lim, ac_r_e_lim)
        ENDIF
        ENDDO
      END SUBROUTINE coeff_bps_small_E
ļ
 Same as coeff_bps_small_E, except returns A-coefficients for a single plasma
  index ib.
ı
      SUBROUTINE a_sing_ib_small_E(nni, ib, ep, zp, mp, betab, zb, mb, nb, ac_s_lim)
      USE physvars
USE mathvars
        IMPLICIT NONE
        INTEGER,
                                       INTENT(IN)
                                                   :: nni
                                                                   number of ions
        INTEGER,
                                       INTENT(IN)
                                                   :: ib
                                                                   plasma species number
        REAL,
                 DIMENSION(1:nni+1), INTENT(IN)
                                                   :: betab
                                                                ļ
                                                                   temp array [1/keV]
        REAL.
                 DIMENSION(1:nni+1), INTENT(IN)
                                                                   mass array [keV]
                                                   :: mb
                                                                ١
                 DIMENSION(1:nni+1),
        REAL,
                                      INTENT(IN)
                                                                   density [1/cc]
                                                   :: nb
                 DIMENSION(1:nni+1),
        REAL,
                                       INTENT(IN)
                                                   :: zb
                                                                ļ
                                                                   charge array
        REAL,
                                       INTENT(IN)
                                                   :: ep
                                                                   projectile energy [ke
        REAL,
                                       INTENT(IN)
                                                   :: mp
                                                                   projectile mass
                                                                                       [ke
        REAL,
                                       INTENT(IN)
                                                                   projectile charge
                                                   :: zp
        REAL,
                                       INTENT(OUT) :: ac_s_lim !
                                                                   A-coeffs [MeV/micron]
        REAL,
                 DIMENSION(1:nni+1)
                                       :: kb2
                                                  [1/cm^2]
                 DIMENSION(1:nni+1)
DIMENSION(1:nni+1)
        REAL,
                                                  [dimensionless]
                                       ::
                                         ab
        REAL,
                                                  [keV]
                                       :: mpb
        REAL,
                 DIMENSION(1:nni+1)
                                               !
                                       :: mbpb
                                                 [dimensionless]
        REAL
                                                 [cm/s]
                 :: vp
                :: vp2
                                                  [cm^2/s^2]
        REAL
                                               ļ
                                                  [1/cm^2]
        REAL
                :: kd2, k2
                   kd, k
        REAL
                                                  [1/cm]
        REAL
                                                  [dimensionless]
                :: a, zp2
        vp = CC*SQRT(2*ep/mp)
                                        [cm/s]
                                        [cm^2/s^2]
        vp2 = vp*vp
        kb2 = DEBYE2*zb*zb*nb*betab
                                        [1/cm^2]
                                        [1/cm^2]
[1/cm]
[1/cm^2]
        kd2 = SUM(kb2)
        kd
           = SQRT(kd2)
            = kb2(1)
        k2
                                                  k2=k_e^2
            = SQRT(k2)
                                                  k = k_e
                                        [1/cm]
        k
        zp2 = zp**2
                                        [dimensionless]
                                        ab=(1/2) betab(ib)*mbc2(ib)*vp2/CC2
        ab =0.5*betab*mb*vp2/CC2
                                        [dimensionless]
        mpb = mp*mb/(mp+mb)
                                      !
                                        [keV]
        mbpb= mb/mpb
                                      ! [dimensionless]
ļ
```

```
a=ab(ib)
                                       ! [dimensionless]
 singular: asymptotic low energy form
        ac_s_{lim}=-(2./3. - 0.4*a)*(LOG(betab(ib)*BEKEV*ABS(zp*zb(ib))* &
          0.5*k*AOCM*mbpb(ib) + 2*GAMMA) - (4./15.)*a
ı
      END SUBROUTINE a_sing_ib_small_E
      SUBROUTINE a_reg_ib_small_E(nni, ib, ep, zp, mp, betab, zb, mb, nb, ac_r_lim)
      USE physvars
USE mathvars
        IMPLICIT NONE
                                        INTENT(IN)
        INTEGER,
                                                      :: nni
                                                                      number of ions
        INTEGER,
                                        INTENT(IN)
                                                      :: ib
                                                                   ١
                                                                      plasma species number
                  DIMENSION(1:nni+1), INTENT(IN)
        REAL,
                                                      :: betab
                                                                   ļ
                                                                      temp array [1/keV]
        REAL,
                                                                   Ţ
                                                                      mass array [keV]
                  DIMENSION(1:nni+1), INTENT(IN)
                                                      :: mb
                  DIMENSION(1:nni+1), INTENT(IN)
        REAL,
                                                                   ļ
                                                      :: nb
                                                                      density [1/cc]
        REAL,
                  DIMENSION(1:nni+1), INTENT(IN)
                                                                      charge array
                                                      :: zb
        REAL,
                                        INTENT(IN)
                                                                   ļ
                                                                      projectile energy [ke
                                                      :: ep
        REAL,
                                        INTENT(IN)
                                                                   İ
                                                                                          [ke
                                                                      projectile mass
                                                      :: mp
        REAL.
                                                                   !
                                        INTENT(IN)
                                                      :: zp
                                                                      projectile charge
        REAL.
                                        INTENT(OUT) :: ac_r_lim !
                                                                      A-coeffs [MeV/micron]
        REAL,
                  DIMENSION(1:nni+1)
                                        :: kb2
                                                    [1/cm^2]
                  DIMENSION(1:nni+1)
DIMENSION(1:nni+1)
DIMENSION(1:nni+1)
        REAL,
                                        :: ab
                                                    [dimensionless]
        REAL,
                                        :: ab2
                                                    [dimensionless]
        REAL,
                                                 ļ
                                                    [keV]
                                        :: mpb
        REAL,
                  DIMENSION(1:nni+1)
                                        :: mbpb !
                                                   [dimensionless]
        REAL
                                                   [cm/s]
                 :: vp
        REAL
                                                 1
                                                   [cm^2/s^2]
                 :: vp2
        REAL
                 :: kd2, k2
                                                   [1/cm^2]
                                                    [1/cm]
[dimensionless]
        REAL
                 :: kd, k
        REAL
                 :: a, zp2
        REAL
                 :: ar1, ar2, c2
                                                 ! [dimensionless]
        vp = CC*SQRT(2*ep/mp)
                                         [cm/s]
                                         [cm^2/s^2]
        vp2 = vp*vp
                                          [1/cm<sup>2</sup>]
[1/cm<sup>2</sup>]
        kb2 = DEBYE2*zb*zb*nb*betab
        kd2 = SUM(kb2)
        kd = SQRT(kd2)
                                          [1/cm]
        k2 = kb2(1)
                                          [1/cm^2]
                                                    k2=k_e^2
                                                    k = k_e
        k
            = SQRT(k2)
                                          [1/cm]
        zp2 = zp**2
                                          [dimensionless]
                                         ab=(1/2) betab(ib)*mbc2(ib)*vp2/CC2
        ab =0.5*betab*mb*vp2/CC2
                                       1
                                         [dimensionless]
        mpb = mp*mb/(mp+mb)
                                       !
                                         [keV]
                                         [dimensionless]
        mbpb= mb/mpb
ļ
        a=ab(ib)
                                          [dimensionless]
        c2=SQRT(a/PI)
                                          [dimensionless]
 regular: asymptotic low energy form
        ab2 = SQRT(ab)
                                          [dimensionless]
                                          [dimensionless] coeff for A_reg with E<<T
        ar1 = -2*SUM(kb2*ab)/k2/5.
        ar2 = SUM(kb2*ab2)/k2
                                         [dimensionless] coeff for A_reg with E<<T
        ar2 = ar2*ar2*PI/30.
        ac_r_{lim}=-c2*((THIRD - 0.2*a)*(LOG(kd2/k2)+1) + ar1 + ar2)
      END SUBROUTINE a_reg_ib_small_E
      SUBROUTINE aq_ib_small_E(nni, ib, ep, zp, mp, betab, zb, mb, nb, aq_lim)
      USE physvars
```

```
USE mathvars
IMPLICIT NONE
        INTEGER,
                                        INTENT(IN)
                                                     :: nni
                                                                     number of ions
        INTEGER,
                                        INTENT(IN)
                                                                     plasma species number
                                                     :: ib
        REAL,
                  DIMENSION(1:nni+1), INTENT(IN)
                                                     :: betab
                                                                     temp array [1/keV]
                  DIMENSION(1:nni+1), INTENT(IN)
        REAL,
                                                     :: mb
                                                                  ļ
                                                                     mass array [keV]
                  DIMENSION(1:nni+1), INTENT(IN)
        REAL,
                                                                  ļ
                                                                     density [1/cc]
                                                     :: nb
        REAL,
                  DIMENSION(1:nni+1),
                                       INTENT(IN)
                                                     :: zb
                                                                  ļ
                                                                     charge array
        REAL,
                                        INTENT(IN)
                                                     :: ep
                                                                  ļ
                                                                     projectile energy [ke
        REAL,
                                        INTENT(IN)
                                                                  ļ
                                                                     projectile mass
                                                                                         [ke
                                                     :: mp
                                                                  !
        REAL,
                                        INTENT(IN)
                                                                     projectile charge
                                                     :: zp
                                                                     A-coeffs [MeV/micron]
        REAL,
                                        INTENT(OUT) :: aq_lim
        REAL,
                  DIMENSION(1:nni+1)
                                        :: kb2
                                                   [1/cm^2]
        REAL,
                  DIMENSION(1:nni+1)
                                                   [dimensionless]
                                        :: ab
        REAL,
                                        :: mpb
                  DIMENSION(1:nni+1)
                                                   [keV]
                  DIMENSION(1:nni+1)
        REAL,
                                        :: mbpb !
                                                  [dimensionless]
        REAL
                                                   [cm/s]
                 :: vp
                                                   [cm^2/s^2]
        REAL
                 :: vp2
                 :: kd2, k2
:: kd, k
                                                   [1/cm^2]
        REAL
                                                   [1/cm]
        REAL
        REAL
                 :: a, zp2
                                                   [dimensionless]
        REAL
                 :: etbar, etbar2
                                                ! [dimensionless]
        vp = CC*SQRT(2*ep/mp)
                                         [cm/s]
        vp2 = vp*vp
                                         [cm^2/s^2]
        kb2 = DEBYE2*zb*zb*nb*betab
                                         [1/cm^2]
        kd2 = SUM(kb2)
                                         [1/cm^2]
        kd = SQRT(kd2)

k2 = kb2(1)
                                         [1/cm]
                                         [1/cm^
                                                   k2=k_e^2
            = SQRT(k2)
                                                   k = k_e
        k
                                         [1/cm]
        zp2 = zp**2
                                         [dimensionless]
                                         ab=(1/2) betab(ib)*mbc2(ib)*vp2/CC2
        ab =0.5*betab*mb*vp2/CC2
                                         [dimensionless]
        mpb = mp*mb/(mp+mb)
                                       1
                                         [keV]
                                         [dimensionless]
        mbpb= mb/mpb
ļ
        a=ab(ib)
                                       ! [dimensionless]
ļ
 quantum: asymptotic low energy form: etbar defined with thermal velocity
ļ
  [\ eta_{pb} = e_p e_b/4\pi \bar \bar v_b $ with \ with \ v_b^2 = 3 T_b/m_b$]
           etbar=4.2115E-3*ABS(zp*zb(ib))*SQRT(betab(ib)*mb(ib))
           etbar2=etbar*etbar
           IF (ib==1) THEN
               aq_lim=LOG(1.5*etbar2)/3. + GAMMA ! electrons only eta_pe << 1
           ELSE
              aq_lim=-1./(27*etbar2)
           ENDIE
      END SUBROUTINE aq_ib_small_E
```

## 3. High Energy Asymptotics

Medium-high energy electrons:

```
acoeff.f90:
!
! ROUTINE: SUBROUTINE coeff_bps_high_E(nni, ep, zp, mp, betab, zb, mb, nb, &
```

```
a_tot_lim, a_i_lim, a_e_lim, ac_tot_lim, ac_i_lim, ac_e_lim, aq_tot_lim,&
    aq_i_lim, aq_e_lim)
 Returns high energy asymptotic regimes. For the ions this means E_p >> T.
! For electrons there are two regimes:
       extreme high energy E_p >> (m_I/m_e)*T [coeff_bps_very_high_E]
 (i)
  (ii) intermediate high energy T << E_p << (m_I/m_e)*T [this routine]
 UNITS: A_b has units of [MeV/micron] (subject to change in updates)
 The incident projectile and the background plasma.
 projectile input quantities:
 ep : classical kinetic energy of the projectile [keV]
 zp : charge of the projectile in units of Z_p [dimensionless]
 mp : mass of the projectile [keV], i.e. mp = mp[grams]*c^2
 plasma input quantities:
      : Number of total plasma species = number ion species + 1
 nni
       : Charges of the plasma species. By convention zp(1) is the
         electron plasma component. [dimensionless, Array]
 betab: Inverse temperatures of the plasma components. For an
         electron-ion plasma, set betab(1)=1/T_e and all other
         values of the array to 1/T_I.
       : Masses of the plasma species [keV].
 mb
 nb
       : Number densities of the plasma species [cm^-3].
 OUTPUT: a_tot_lim, a_i_lim, a_e_lim, ac_tot_lim, ac_i+lim, ac_e_lim,
          aq_tot_lim, aq_i_lim, aq_e_lim
 classical electron
                       : ac_e_lim
                        : ac_i_lim [sum over all ions]
! classical ion
! classical total
                        : ac_tot_lim = ac_e_lim + ac_i_lim
 quantum
             electron
                       : aq_e_lim
                          aq_i_lim [sum over all ions]
a_tot_lim = aq_e_lim + aq_i_lim
 quantum
             ion
 quantum
             total
                        : a_e_lim = ac_e_lim + aq_e_lim
! total
             electric
                          a_i_lim = ac_i_lim + aq_i_lim
! total
 total
                        ! a_{tot_lim} = a_{e_lim} + a_{i_lim}
      SUBROUTINE coeff_bps_high_E(nni, ep, zp, mp, betab, zb, mb, nb, a_tot_lim, a_i_lim, a_e_lim, ac_tot_lim, ac_i_lim, ac_e_lim, aq_tot_lim, aq_i_lim, aq_e_lim, ac_s_i_lim, ac_s_e_lim,
             ac_r_i_lim, ac_r_e_lim)
      USE physvars
USE mathvars
        IMPLICIT NONE
                                                      :: nni
        INTEGER,
                                        INTENT(IN)
                                                                    number of ions
        REAL,
                  DIMENSION(1:nni+1), INTENT(IN)
                                                      :: betab
                                                                                    [1/keV]
                                                                    temp array
        REAL,
                  DIMENSION(1:nni+1), INTENT(IN)
                                                      :: mb
                                                                    mass array
                                                                                    [keV]
        REAL,
                  DIMENSION(1:nni+1), INTENT(IN)
                                                      :: nb
                                                                    density array [1/cc]
                  DIMENSION(1:nni+1), INTENT(IN)
                                                                    charge array
        REAL,
                                                      :: zb
                                                                   Projectile
        REAL,
                                        INTENT(IN)
                                                      :: ep
                                                                    projectile energy [keV]
        REAL,
                                        INTENT(IN)
                                                      :: mp
                                                                    projectile mass
                                                                                         [keV]
        REAL,
                                        INTENT(IN)
                                                      :: zp
                                                                    projectile charge
                                                                      ! A-coeffs [MeV/micron
                                        INTENT(OUT) :: a_tot_lim
        REAL,
                                                                         electron + ion
                                        INTENT(OUT) :: a_i_lim
        REAL,
                                                                         ion contribution
        REAL,
                                        INTENT(OUT) :: a_e_lim
                                                                         electron contributi
                                        INTENT(OUT) :: ac_tot_lim !
INTENT(OUT) :: ac_i_lim !
        REAL,
                                                                         classical
        REAL,
                                                                         classical
```

```
INTENT(OUT) :: ac_e_lim !
INTENT(OUT) :: aq_tot_lim !
         REAL,
                                                                            classical
         REAL,
                                                                            quantum
                                          INTENT(OUT) :: aq_i_lim
         REAL,
                                                                            quantum
         REAL,
                                          INTENT(OUT) :: aq_e_lim
                                                                            quantum
                                          INTENT(OUT) :: ac_s_i_lim
INTENT(OUT) :: ac_s_e_lim
INTENT(OUT) :: ac_r_i_lim
         REAL,
         REAL,
         REAL,
         REAL,
                                          INTENT(OUT) :: ac_r_e_lim
        REAL :: ac_r_lim, ac_s_lim, aq_lim INTEGER :: ib, nnb
                  :: vp
         REAL
                                                      [cm/s]
         REAL
                  :: vp2
                                                      [cm^2/s^2]
         REAL
                  :: a, zp2
                                                      [dimensionless]
         REAL
                  :: c1, cs
                                                       [keV/cm]
         REAL
                  :: c2
                                                       [dimensionless]
                                                       [1/cm]
         REAL
                  :: k
                  :: k2, ke2
                                                       [1/cm^2]
         REAL
         R.E.A.L.
                  :: eta
                                                       [dimensionless]
                                                       [keV]
         REAL
                  :: te, mec2, mpec22
                   DIMENSION(1:nni+1)
         REAL,
                                          :: ab
                                                       [dimensionless]
                   DIMENSION(1:nni+1)
                                                    ! [keV]
                                          :: mpb
         REAL,
                   DIMENSION(1:nni+1)
                                          :: kb2
         REAL.
                                                    ! [1/cm<sup>2</sup>
 omi2, and omi needed only for
 E >> (mI/me)*T: extreme high energy limit for electrons
                  :: omi2, omi
ļ
         REAL
                                                    ! [1/s^2, 1/s]]
         nnb = nni+1
         vp = CC*SQRT(2*ep/mp)
                                            [cm/s]
                                            [cm^2/s^2]
         vp2 = vp*vp
                                            [1/cm<sup>2</sup>]
[1/cm<sup>2</sup>]
         kb2 = DEBYE2*zb*zb*nb*betab
         k2 = kb2(1)
                                                       k2=k_e^2
            = SQRT(k2)
                                                       k = k_e
                                            [1/cm]
         zp2 = zp**2
                                            [dimensionless]
                                         ! ab=(1/2) betab(ib)*mb(ib)*vp2/CC2
         ab = 0.5*betab*mb*vp2/CC2
                                         ! [dimensionless]
         mpb = mp*mb/(mp+mb)
                                         ! [keV]
  initialize A-coefficients
         a_{tot_lim} = 0
                         ! electron + ion
                  = 0
                         ! ion contribution
         a_i_lim
                    = 0
                          ! electron contribution
         a_e_lim
         ac_tot_lim= 0
                            classical total
                          ! classical electron
         ac_e_lim = 0
         ac_ilim = 0
                          ! classical ion
         aq_tot_lim= 0
                          ! quantum total
         aq_e_lim = 0

aq_i_lim = 0
                          ! quantum electron
                          ! quantum ion
         ac_s_i_lim=0
         ac_s_e_lim=0
         ac_r_i_lim=0
ac_r_e_lim=0
١
         DO ib=1,nni+1
         IF (zb(ib) .NE. O.) THEN
            a=ab(ib)
                                             ! [dimensionless]
            c1=2*zp2*BEKEV*kb2(ib)*AOCM ! [keV/cm]
            c1=c1*1.E-7
                                               [MeV/micron]
            c2=SQRT(a/PI)
                                               [dimensionless]
 singular: asymptotic high energy form [need electrons]
```

```
At this point I only have an asymptotic form for the total
  electron contribution (classical + quantum = sing + reg + quantum). I'll write this expression to ac_s_lim for now. This will give aq_lim=0 and ac_lim=a_e. FIX LATER.
            IF (ib == 1) THEN
  T \ll E \ll (mI/me)*T: intermediate high energy limit for electrons
                te=1./betab(ib)
                mpec22=mpb(ib)**2
                mec2 = mb(ib)
                       =kb2(ib)
                ke2
                ac_s_lim=THIRD*(LOG(8*te*mpec22/(mec2*HBARC**2*ke2)) - GAMMA -1)
                                                           ! [MeV/micron]
                ac_s_lim=c1*c2*ac_s_lim
            ELSE
                cs=0.5*c1/ab(ib)
                ac_s_lim=(LOG(ABS(zp*zb(ib))*BEKEV*& ! [dimensionless]
                  k*AOCM*CC2/(mpb(ib)*vp2)) + GAMMA) !
                ac_s_lim=-cs*ac_s_lim
                                                             [MeV/micron]
            ENDIF
  regular: asymptotic high energy form [need electrons]
            IF (ib == 1) THEN
                ac_r_lim=0
            ELSE
                 ac_r_{lim}=-0.25*c1/ab(ib)
                                                                        ! [MeV/micron]
            ENDIF
  quantum: asymptotic high energy form [need electrons]
            IF (ib == 1) THEN
                aq_lim=0
            ELSE
                eta =ABS(zp*zb(ib))*2.1870E8/vp ! [dimensionless] quantum parameter
                aq_lim=LOG(eta) + GAMMA
                aq_lim=aq_lim*c1/a/2
            ENDIF
            CALL a_collect(ib,nnb,ac_s_lim,ac_r_lim,aq_lim,a_tot_lim,a_i_lim,
               a_e_lim,ac_tot_lim,ac_i_lim,ac_e_lim,aq_tot_lim,aq_i_lim,aq_e_lim, &
               ac_s_i_lim, ac_s_e_lim, ac_r_i_lim, ac_r_e_lim)
         ENDIF
         ENDDO
       END SUBROUTINE coeff_bps_high_E
Extreme-high energy electrons:
acoeff.f90:
       SUBROUTINE coeff_bps_very_high_E(nni, ep, zp, mp, betab, zb, mb, nb, a_tot_lim, a_i_lim, a_e_lim, ac_tot_lim, ac_i_lim, ac_e_lim,
                                                                                      &
             aq_tot_lim, aq_i_lim, aq_e_lim, ac_s_i_lim, ac_s_e_lim,
             ac_r_i_lim, ac_r_e_lim)
      USE physvars USE mathvars
         IMPLICIT NONE
                   DIMENSION(1:nni+1), INTENT(IN)
         INTEGER,
                                                                       number of ions
                                                        :: nni
                                                                                        [1/keV]
         REAL,
                                                        :: betab
                                                                       temp array
                   DIMENSION(1:nni+1), INTENT(IN)
         REAL,
                                                                    ļ
                                                        :: mb
                                                                                        [keV]
                                                                       mass array
         REAL,
                   DIMENSION(1:nni+1), INTENT(IN)
                                                        :: nb
                                                                       density array [1/cc]
         REAL,
                   DIMENSION(1:nni+1), INTENT(IN)
                                                        :: zb
                                                                       charge array
```

```
Projectile
       REAL,
                                         INTENT(IN)
                                                                      projectile energy [keV]
                                                       :: ep
       REAL,
                                         INTENT(IN)
                                                       :: mp
                                                                      projectile mass
       REAL,
                                         INTENT(IN)
                                                                      projectile charge
                                                       :: zp
                                                                        ! A-coeffs [MeV/micron
       REAL,
                                                                           electron + ion
                                         INTENT (OUT)
                                                       :: a_tot_lim
                                         INTENT (OUT)
INTENT (OUT)
INTENT (OUT)
       REAL,
                                                       :: a_i_lim
                                                                           ion contribution
       REAL,
                                         INTENT(OUT) :: a_e_lim
INTENT(OUT) :: ac_tot_lim
INTENT(OUT) :: ac_i_lim
                                                                           electron contributi
       REAL,
                                                                           classical
       REAL,
                                                                           classical
                                         INTENT (OUT)
       REAL,
                                                      :: ac_e_lim
                                                                           classical
       REAL,
                                         INTENT(OUT) :: aq_tot_lim !
                                                                           quantum
                                         INTENT(OUT) :: aq_i_lim
       REAL,
                                                                           quantum
       REAL,
                                         INTENT(OUT) :: aq_e_lim
                                                                           quantum
       REAL,
                                         INTENT(OUT) :: ac_s_i_lim
                                         INTENT(OUT)
INTENT(OUT)
       REAL,
                                                      :: ac_s_e_lim
       REAL,
                                                      :: ac_r_i_lim
                                         INTENT(OUT) :: ac_r_e_lim
       REAL,
                :: omi2, omi
:: ac_r_lim, ac_s_lim, aq_lim
       REAL
       REAL
       INTEGER :: ib, nnb
                :: vp
                                                     [cm/s]
       REAL
       REAL
                                                     [cm^2/s^2]
                :: vp2
       REAL
                :: a, zp2
                                                     [dimensionless]
       REAL
                :: c1, cs
                                                     [keV/cm]
       REAL
                :: c2
                                                     [dimensionless]
       REAL
                :: k
                                                     [1/cm]
                :: k2
                                                     [1/cm^2]
       REAL
                                                     [dimensionless]
       REAL
                :: eta
       REAL,
                 DIMENSION(1:nni+1)
                                         :: ab
                                                     [dimensionless]
                 DIMENSION(1:nni+1)
                                                     [keV]
                                         :: mpb
       REAL,
                                                     [1/cm^2]
       REAL,
                 DIMENSION(1:nni+1)
                                         :: kb2
omi2, and omi needed only for
E >> (mI/me)*T: extreme high energy limit for electrons
       REAL
                                                   ! [1/s^2, 1/s]]
                :: omi2, omi
       nnb = nni+1
       vp = CC*SQRT(2*ep/mp)
                                          [cm/s]
       vp2 = vp*vp
                                          [cm^2/s^2]
                                          [1/cm<sup>2</sup>]
[1/cm<sup>2</sup>]
       kb2 = DEBYE2*zb*zb*nb*betab
       k2 = kb2(1)
                                                      k2=k_e^2
           = SQRT(k2)
                                           [1/cm]
                                                      k = k_e
                                          [dimensionless]
       zp2 = zp**2
                                        ! ab=(1/2) betab(ib)*mb(ib)*vp2/CC2
                                        ! [dimensionless]
       ab = 0.5*betab*mb*vp2/CC2
                                        ! [keV]
       mpb = mp*mb/(mp+mb)
initialize A-coefficients
       a_{tot_lim} = 0
                         ! electron + ion
       a_i_lim
                  = 0
                        ! ion contribution
                  = 0
                         ! electron contribution
       a_e_lim
       ac_tot_lim= 0
ac_e_lim = 0
                         ! classical total
! classical electron
                 = 0
       ac_i_lim
                         ! classical ion
       aq_tot_lim= 0
                        ! quantum total
       aq_e_lim = 0

aq_i_lim = 0
                         ! quantum electron
                         ! quantum ion
       ac_s_i_{lim=0}
       ac_s_e_lim=0
```

[keV]

```
ac_r_i_lim=0
         ac_r_e_lim=0
ļ
         DO ib=1,nni+1
         IF (zb(ib) .NE. O.) THEN
            a=ab(ib)
                                                 [dimensionless]
             c1=2*zp2*BEKEV*kb2(ib)*AOCM ! [keV/cm]
            c1=c1*1.E-7
                                                 [MeV/micron]
             c2=SQRT(a/PI)
                                                 [dimensionless]
  singular: asymptotic high energy form [need electrons]
ļ
 At this point I only have an asymptotic form for the total
 electron contribution (classical + quantum = sing + reg + quantum). I'll write this expression to ac_s_lim for now. This will give
  aq_lim=0 and ac_lim=a_e. FIX LATER.
            IF (ib == 1) THEN
ı
 E >> (mI/me)*T: extreme high energy limit for electrons
                omi2=OMEGI2*zb(ib)*zb(ib)*nb(ib)*AMUKEV/mb(ib) ! [1/s^2]
                omi =SQRT(omi2)
                ac_s_lim=0.5*c1*LOG(2*mpb(ib)*vp2/(HBARC*CC*omi))/ab(ib)
            ELSE
                cs=0.5*c1/ab(ib)
                ac_s_lim=(LOG(ABS(zp*zb(ib))*BEKEV*& ! [dimensionless]
                   k*AOCM*CC2/(mpb(ib)*vp2)) + GAMMA) !
                ac_s_lim=-cs*ac_s_lim
                                                               [MeV/micron]
            ENDIF
ļ
 regular: asymptotic high energy form [need electrons]
             IF (ib == 1) THEN
                ac_r_lim=0
            ELSE
                                                                           ! [MeV/micron]
                  ac_r_{lim}=-0.25*c1/ab(ib)
            ENDIF
  quantum: asymptotic high energy form [need electrons]
             IF (ib == 1) THEN
                aq_lim=0
                eta =ABS(zp*zb(ib))*2.1870E8/vp ! [dimensionless] quantum parameter
                aq_lim=LOG(eta) + GAMMA
                aq_lim=aq_lim*c1/a/2
            ENDIF
             \begin{array}{llll} \texttt{CALL} & \texttt{a\_collect(ib,nnb,ac\_s\_lim,ac\_r\_lim,aq\_lim,a\_tot\_lim,a\_i\_lim,} & \texttt{a\_e\_lim,ac\_tot\_lim,ac\_i\_lim,ac\_e\_lim,aq\_tot\_lim,aq\_i\_lim,aq\_e\_lim,} \\ \end{array} 
               ac_s_i_lim, ac_s_e_lim, ac_r_i_lim, ac_r_e_lim)
         ENDIF
         ENDDO
       END SUBROUTINE coeff_bps_very_high_E
```