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C-Coefficient in Clog

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Abstract

Physics documentation for the BPS stopping power in the code Clog.

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I. GENERAL ANALYTIC EXPRESSIONS FOR THE BPS C-COEFFICIENTS

Suppose we have a plasma with various species labeled by an index b at distinct temperatures T_b , number densities n_b , and species masses m_b .¹ Temperature will be measured in energy units, and we denote the inverse temperature by $\beta_b = 1/T_b$. Electrostatic units will be rationalized cgs. The projectile will have mass m_a , charge e_a and energy $E_a = \frac{1}{2} m_a v_a^2$. The BPS $\mathcal{C}^{\ell\ell}$ -coefficients take the form

$$C_{ab}^{\ell\ell} = \left(C_{ab,R}^{\ell\ell <} + C_{ab,S}^{\ell\ell C}\right) + C_{ab}^{\ell\ell \Delta Q} \tag{1.1}$$

with

$$C_{ab,R}^{\ell\ell<} = \frac{e_a^2}{4\pi} \frac{1}{\beta_b v_a} \frac{i}{2\pi} \int_{-1}^1 \frac{d\cos\theta}{\cos\theta} \frac{\rho_b(v_a\cos\theta)}{\rho_{\text{total}}(v_a\cos\theta)} F(v_a\cos\theta) \ln\left\{\frac{F(v_a\cos\theta)}{K^2}\right\}, \quad (1.2)$$

$$C_{ab,s}^{\ell\ell C} = \frac{e_a^2 \kappa_b^2}{4\pi} \left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} \frac{1}{\beta_b} \int_0^1 du \, u^{-1/2} \exp\left\{-\frac{1}{2} \beta_b m_b v_a^2 u\right\} \left[-\ln\left(\beta_b \frac{e_a e_b}{4\pi} K \frac{m_b}{m_{ab}} \frac{u}{1-u}\right) - 2\gamma\right]$$
(1.3)

$$C_{ab}^{\ell\ell} \Delta Q = -\frac{e_a^2 \kappa_b^2}{4\pi} \left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} \frac{1}{\beta_b v_a} \int_0^\infty dv_{ab} \left\{ \text{Re} \, \psi \left(1 + i\eta_{ab}\right) - \ln \eta_{ab} \right\}$$

$$\left[\exp \left\{ -\frac{1}{2} \beta_b m_b \left(v_a - v_{ab}\right)^2 \right\} - \exp \left\{ -\frac{1}{2} \beta_b m_b \left(v_a + v_{ab}\right)^2 \right\} \right], \tag{1.4}$$

This regular form is BPS (9.7); the singular form is BPS (9.5); the quantum form is BPS (10.27), where

$$\eta_{ab} = \frac{e_a e_b}{4\pi \hbar v_{ab}} \ . \tag{1.5}$$

The Debye wavenumber K is arbitrary and will typically be chosen as $K = \kappa_e$. The function F(v) takes the form

$$F(v) = -\int_{-\infty}^{\infty} du \, \frac{\rho_{\text{tot}}(u)}{v - u + i\eta} \quad \text{with} \quad \rho_{\text{tot}}(u) = \sum_{b} \rho_{b}(u)$$
 (1.6)

$$\rho_b(v) = \kappa_b^2 \sqrt{\frac{\beta_b m_b}{2\pi}} v \exp\left\{-\frac{1}{2} \beta_b m_b v^2\right\} , \qquad (1.7)$$

and its relation to the dielectric function is

$$k^{2} \epsilon(\mathbf{k}, \mathbf{k} \cdot \mathbf{v}) = k^{2} + F(\hat{\mathbf{k}} \cdot \mathbf{v}) . \tag{1.8}$$

¹ By convention b = 1 will be the electron component.

The first term $C_{ab,\mathrm{R}}^{\ell\ell}$ arises from long-distance collective effects from the dielectric function, and it involves all plasma species (even species c different from a and b). This is the term I call non-separable, meaning that it cannot be written as a sum of individual plasma components involving only a single species. The second term $C_{ab,\mathrm{S}}^{\ell\ell}$ arises from short-distance two-body classical scattering, and the third term $C_{ab}^{\ell\ell\,\mathrm{QM}}$ is the two-body quantum scattering correction to all orders in the quantum parameters $\bar{\eta}_{ab}$. Three body and higher effects are contained in our systematic error term, the next-to-next-to-leading order term proportional to g^3 . In a strongly coupled plasma these higher order effects dominate, but in a weakly coupled plasma they are negligible.

II. THE MAIN DRIVER

I will return the C-coefficients in three forms:

- i. bps_ccoeff_ab_mass: For a given pair of indices p and b (the projectile p will often be denoted by species index a), this routine returns the individual component $C_{ab}^{\ell\ell}(E)$ for a given energy E. The quantum parameter η can be arbitrary. This routine is used to construct the entries in the next two subroutines.
- ii. bps_ccoeff_ab_matrix: Returns the complete matrix of coefficients $C_{ab}^{\ell\ell}(E)$.
- iii. bps_ccoeff_ei_mass: This routine returns the sum over the ions $C_{pi}^{\ell\ell} = \sum_i C_{pi}^{\ell\ell}$ for a given projectile p. It also returns the coulomb logarithm.

A. The Driver Routine: bps_ccoeff_ab_mass

This subroutine returns the matrix of values $C_{ab}^{\ell\ell}(E)$ for a given energy E. The driving routine that calls and assembles the singular, regular, and quantum pieces.

ccoeff.f90:bps_ccoeff_ab_mass

```
!*** still need to fix scaling/units ***
! main driver for C-coefficient for general quantum and electron-mass regimes
SUBROUTINE bps_ccoeff_ab_mass(nni, ep, mp, zp, ia, ib, betab, zb, mb, nb, &
            c_ab, c_ab_sing, c_ab_reg, c_ab_qm)
     USE physvars
USE mathvars
        IMPLICIT NONE
                                                                     Plasma:
       INTEGER,
                                             INTENT(IN)
                                                         :: nni
                                                                      number of ions
       REAL,
                                             INTENT(IN)
                                                         :: ep
                                                                       energy input [ke
       REAL,
                                             INTENT(IN)
                                                                      mass [keV]
                                                         :: mp
       REAL,
                                             INTENT(IN)
                                                         :: zp
                                                                       charge
       INTEGER,
                                             INTENT(IN)
                                                         :: 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
                 DIMENSION(1:nni+1),
       REAL,
                                             INTENT(IN)
                                                                      density [1/cc]
                                                         :: nb
                 DIMENSION(1:nni+1),
       REAL,
                                             INTENT(IN)
                                                         :: zb
                                                                       charge array
                                                                     C-coeffs [MeV/mic
       REAL,
                                             INTENT(OUT) :: c_ab
                                             INTENT(OUT) :: c_ab_sing
       REAL,
       REAL,
                                             INTENT(OUT) :: c_ab_reg
       REAL,
                                             INTENT(OUT) :: c_ab_qm
                                     :: mpb, mbpb, kb2, ab
:: vp, zp2, k, k2, kd, kd2, a, b, eta
:: cc_r, cc_s, cq, c1, c2, c3
       REAL,
                 DIMENSION(1:nni+1)
       REAL
       REAL
                                     :: EPS_SMALL_E=2.E-4
:: EPS_SMALL_E_SING=2.E-4
:: EPS_SMALL_E_REG=2.E-4
       REAL, PARAMETER
       REAL, PARAMETER REAL, PARAMETER
 initialize components of C-coefficients
```

kb2=8*PI*AOCM*BEKEV*zb*zb*nb*betab

```
[1/cm<sup>2</sup>]
[1/cm]
[1/cm<sup>2</sup>]
        kd2 = SUM(kb2)
kd = SQRT(kd2)
k2 = kb2(1)
            = SQRT(k2)
                                           [1/cm]
                                                    k = k_e
 Loop over charged plasma species
        mpb = mp*mb/(mp+mb)
                                          ! [keV]
        mbpb= mb/mpb
                                          ! [dimensionless]
        vp = CC*SQRT(2*ep/mp)
                                           [cm/s]
        zp2 = zp**2
                                           [dimensionless]
        ab = 0.5*betab*mb*vp*vp/CC2
                                          ! [dimensionless]
                                          ! ab=(1/2) betab(ib)*mbc2(ib)*vp2/CC2
        IF (zb(ib) .NE. O.) THEN
               =ab(ib)
              =-Log(2*betab(ib)*BEKEV*ABS(zp*zb(ib))*k*AOCM*mbpb(ib) )-2*GAMMA
            eta=ABS(zp*zb(ib))*2.1870E8/vp ! defined with projectile velocity vp
                                               [keV/cm] c1 = e_p^2 \text{ kappa_b^2/(4 Pi)}
            c1=2*zp2*BEKEV*kb2(ib)*AOCM
            c1=c1*1.E-7
                                               [MeV/micron]
           c2=SQRT(a/PI)
                                               [dimensionless] c2=SQRT(betab(ib)*mb(ib)/
           C3=CC/(betab(ib)*vp)
                                   ! 1/betab(ib)*vp note: dE_per/dx = C/m*c^2
            c3=c3/1000.
                                    ! convert from KeV to MeV
  C_{ab}-classical-singular
        CALL c_sing_mass(a,b,cc_s)
        c_ab_sing=c1*c2*c3*cc_s
  C_{ab}-classical-regular
        CALL c_reg_mass(nni,ia,ib,vp,k2,kb2,betab,mb,cc_r)
        c_ab_reg=c1*c3*cc_r
  C_{ab}-quantum
        CALL c_quantum_mass(ia,ib,a,eta,cq) ! eta = dimensionless quantum param.
        c_ab_qm=c1*c2*c3*cq
  C_{ab}-total
        c_ab=c_ab_sing + c_ab_reg + c_ab_qm
        ENDIF
      END SUBROUTINE bps_ccoeff_ab_mass
ccoeff.f90:bps_ccoeff_ab_matrix
SUBROUTINE bps_ccoeff_ab_matrix(nni, ep, betab, zb, mb, nb, &
  c_ab, c_ab_sing, c_ab_reg, c_ab_qm, c_tot, c_i, c_e, cc_tot, &
        cc_i, cc_e, cq_tot, cq_i, cq_e, cc_s_i, cc_s_e, cc_r_i, cc_r_e)
      USE physvars
USE mathvars
                                                                        Plasma:
        IMPLICIT NONE
                                               INTENT(IN)
                                                            :: nni
        INTEGER,
                                                                          number of ions
                                               INTENT(IN)
        REAL,
                                                            :: 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]
                  DIMENSION(1:nni+1),
                                               INTENT(IN)
        REAL,
                                                            :: nb
                                                                          density [1/cc]
                                                                       ! C-coeffs [MeV/mic
```

```
REAL,
  REAL,
  REAL,
            DIMENSION(1:nni+1,1:nni+1),INTENT(OUT) :: c_ab_reg
  REAL,
            DIMENSION(1:nni+1,1:nni+1),INTENT(OUT) :: c_ab_qm
  REAL,
                                            INTENT(OUT)
INTENT(OUT)
            DIMENSION(1:nni+1),
                                                          :: c_tot
            DIMENSION(1:nni+1),
DIMENSION(1:nni+1),
  REAL,
                                                          :: c_i
:: c_e
  REAL,
                                            INTENT (OUT)
  REAL,
            DIMENSION(1:nni+1),
                                            INTENT (OUT)
                                                          :: cc_tot
            DIMENSION (1:nni+1),
                                            INTENT (OUT)
INTENT (OUT)
INTENT (OUT)
  REAL,
                                                          :: cc_i
  REAL,
            DIMENSION(1:nni+1),
DIMENSION(1:nni+1),
                                                         :: cc_e
:: cq_tot
  REAL,
                                            INTENT(OUT) :: cq_i
  REAL,
            DIMENSION(1:nni+1),
                                            INTENT(OUT) :: cq_e
  REAL.
            DIMENSION(1:nni+1),
  REAL,
            DIMENSION(1:nni+1),
                                            INTENT(OUT)
                                                         :: cc_s_i
  REAL,
            DIMENSION(1:nni+1),
DIMENSION(1:nni+1),
                                            INTENT(OUT)
INTENT(OUT)
                                                         :: cc_s_e
:: cc_r_i
  REAL,
            DIMENSION(1:nni+1),
                                            INTENT(OUT) :: cc_r_e
  REAL.
  R.E.A.L.
           :: cab, cab_sing, cab_reg, cab_qm
  REAL :: mp, zp
INTEGER :: ia, ib
  c_i
  cc_s_i = 0
  cc_r_i = 0
  cc_i = 0
  cq_i = 0
D0 ia=1,nni+1
mp=mb(ia)
    zp=zb(ia)
    DD ib=1,nni+1
      CALL bps_ccoeff_ab_mass(nni, ep, mp, zp, ia, ib, betab, zb, mb, nb, &
       cab, cab_sing, cab_reg, cab_qm) !*! change to bps_acoeff_ab_mass
       c_ab(ia,ib)
                         =cab
       c_ab_sing(ia,ib)=cab_sing
       c_ab_reg(ia,ib) =cab_reg
                        =cab_qm
       c_ab_qm(ia,ib)
       IF (ib == 1) THEN
          c_e(ia)
                     = cab
          cc_s_e(ia) = cab_sing
          cc_r_e(ia)= cab_reg
          cc_e(ia) = cab_sing + cab_reg
          cq_e(ia)
                     = cab_qm
       ELSE
          c_i(ia)
                      = c_i(ia)
                                     + cab
          cc_s_i(ia) = cc_s_i(ia) + cab_sing
          cc_r_i(ia) = cc_r_i(ia) + cab_reg
          cc_i(ia)
                     = cc_i(ia)
                                     + cab_sing + cab_reg
          cq_i(ia) = cq_i(ia)
                                     + cab_qm
       ENDIF
    ENDDO
    c_{tot}(ia) = c_{e}(ia) + c_{i}(ia)

c_{tot}(ia) = c_{e}(ia) + c_{i}(ia)
    cq_tot(ia) = cq_e(ia) + cq_i(ia)
  ENDDO
END SUBROUTINE bps_ccoeff_ab_matrix
```

ccoeff.f90:bps_ccoeff_ei_mass

```
SUBROUTINE bps_ccoeff_ei_mass(nni, ep, zp, mp, betab, zb, mb, nb, & c_tot, c_i, c_e, cc_tot, cc_i, cc_e, cq_tot, cq_i, cq_e, &
            cc_s_i, cc_s_e, cc_r_i, cc_r_e)
      USE physvars
      USE mathvars
      USE controlvars
        IMPLICIT NONE
                                                              ! Plasma:
        INTEGER,
                                                                 number of ions
                                      INTENT(IN)
                                                   :: nni
        REAL,
                 DIMENSION(1:nni+1), INTENT(IN)
                                                   :: betab
                                                                 temp array [1/keV]
        REAL,
                 DIMENSION(1:nni+1), INTENT(IN)
                                                   :: mb
                                                                 mass array [keV]
                                                                 density [1/cc]
        REAL,
                 DIMENSION(1:nni+1), INTENT(IN)
                                                   :: nb
                 DIMENSION(1:nni+1), INTENT(IN)
        REAL,
                                                                 charge array
                                                   :: zb
                                                                Projectile
        REAL,
                                      INTENT(IN)
                                                                 projectile energy [keV]
                                                   :: ep
        REAL,
                                      INTENT(IN)
                                                   :: mp
                                                                 projectile mass
                                                                                    [keV]
        REAL,
                                      INTENT(IN)
                                                   :: zp
                                                                 projectile charge
                                                                C-coeffs [MeV/micron]
                                      INTENT(OUT) :: c_tot
                                                                 electron + ion
        REAL,
                                      INTENT(OUT) :: c_i
INTENT(OUT) :: c_e
        REAL,
                                                                 ion contribution
        REAL,
                                                  :: c_e
                                                                 electron contribution
        REAL,
                                      INTENT(OUT)
                                                  :: cc_tot
                                                                 classical
                                      INTENT(OUT)
INTENT(OUT)
        REAL,
                                                   :: cc_i
                                                                 classical
        REAL,
                                                   :: cc_e
                                                                 classical
                                      INTENT(OUT) :: cq_tot
        REAL,
                                                                 quantum
                                      INTENT(OUT) :: cq_i
        REAL,
                                                                 quantum
                                      INTENT(OUT) :: cq_e
        REAL,
                                                                 quantum
        REAL,
                                      INTENT (OUT)
                                                   :: cc_s_i
                                      INTENT(OUT) :: cc_s_e
INTENT(OUT) :: cc_r_i
INTENT(OUT) :: cc_r_e
        REAL,
        REAL,
        REAL,
                 :: cdum, cc_s, cc_r, cq
        INTEGER :: ia, ib, nnb
ı
 initialize components of C-coefficients
        c_{tot} = 0
                  ! electron + ion
        c_i
              =0
                   ! ion contribution
                  ļ
              =0
                    electron contribution
        c_e
        cc_tot=0
                   ! classical total
        cc_e =0
cc_i =0
                   ! classical electron ! classical ion
        cq_tot=0
                   ! quantum total
        cq_e =0
cq_i =0
                   ! quantum electron
        cq_i
                   ! quantum ion
        c\bar{c}_s=0
        cc_s_e=0
        cc_r_i=0
        cc_r_e=0
        NNB = nni+1
                                      ! number of ions + electrons
        ia=1
        DO ib=1,nni+1
          (zb(ib) .NE. O.) THEN
            CALL bps_ccoeff_ab_mass(nni, ep, mp, zp, ia, ib, betab, zb, mb, nb, &
            cdum, cc_s, cc_r, cq)
            CALL x_collect(ib, NNB, cc_s, cc_r, cq,
```

B. The Regular Contribution: c_reg_mass

The long-distance regular contribution can be expressed as

$$C_{ab,R}^{\ell\ell} = \frac{e_a^2}{4\pi} \frac{1}{\beta_b v_a} \frac{i}{2\pi} \int_{-1}^1 \frac{du}{u} \frac{\rho_b(v_a u)}{\rho_{\text{total}}(v_a u)} F(v_a u) \ln \left\{ \frac{F(v_a u)}{K^2} \right\}$$

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

$$= -\frac{e_a^2}{4\pi} \frac{1}{\beta_b v_a} \frac{1}{2\pi} \int_0^1 \frac{du}{u} \frac{\rho_b(v_a u)}{\rho_{\text{total}}(v_a u)} H(v_a u) ,$$

$$(2.3)$$

where we have defined

$$H(v) \equiv -i \left[F(v) \ln \left\{ \frac{F(v)}{K^2} \right\} - F^*(v) \ln \left\{ \frac{F^*(v)}{K^2} \right\} \right] = 2 \left[F_{\text{Re}} \arg\{F\} + F_{\text{Im}} \ln \left\{ \frac{|F|}{K^2} \right\} \right]. \tag{2.4}$$

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

$$F(v) = K^2 \mathbb{F}(v) \text{ and } H(v) = K^2 \mathbb{H}(v) .$$
 (2.5)

Defining the parameters

$$a_c \equiv \left(\frac{\beta_c m_c}{2}\right)^{1/2} \tag{2.6}$$

$$\bar{\kappa}_c^2 \equiv \frac{\kappa_c^2}{K^2} \tag{2.7}$$

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

$$\mathbb{F}_{\text{Re}}(\{a_c v\}, \{\bar{\kappa}_c\}) = \sum_c \bar{\kappa}_c^2 \left(1 - 2a_c v \operatorname{daw}\{a_c v\}\right)$$
(2.8)

$$\mathbb{F}_{\text{Im}}(\{a_c v\}, \{\bar{\kappa}_c\}) = \sqrt{\pi} \sum_c \bar{\kappa}_c^2 a_c v e^{-a_c^2 v^2}.$$
 (2.9)

The ratio of weighting factors can be written in terms of a function \mathbb{R}_{ab} defined by

$$\frac{\rho_b(v_a u)}{\rho_{\text{total}}(v_a u)} \cdot H(v_a u) = K^2 \frac{\rho_b(v_a u)}{\rho_{\text{total}}(v_a u)} \cdot \mathbb{H}(v_a u)$$
(2.10)

$$= K^{2} \frac{\kappa_{b}^{2} (\beta_{b} m_{b}/2\pi)^{1/2} v_{a} u e^{-\frac{1}{2} \beta_{b} m_{b} v_{a}^{2} u^{2}}}{\sum_{c} \kappa_{c}^{2} (\beta_{c} m_{c}/2\pi)^{1/2} v_{a} u e^{-\frac{1}{2} \beta_{c} m_{c} v_{a}^{2} u^{2}}} \cdot \mathbb{H}(v_{a} u)$$
(2.11)

$$= \kappa_b^2 \cdot \underbrace{\left[\sum_c \frac{\kappa_c^2}{K^2} \left(\frac{\beta_c m_c}{\beta_b m_b}\right)^{1/2} e^{\frac{1}{2}(\beta_b m_b - \beta_c m_c) v_a^2 u^2}\right]^{-1}}_{\mathbb{R}_{ab}(v_a u)} \cdot \mathbb{H}(v_a u) (2.12)$$

$$= \kappa_b^2 \,\mathbb{R}_{ab}(v_a u) \,\mathbb{H}(v_a u) \ . \tag{2.13}$$

We can now express the regular piece as

$$C_{ab,R}^{\ell\ell C} = \underbrace{\left[\frac{e_a^2 \kappa_b^2}{4\pi}\right]}_{C_{ab,R}} \cdot \frac{1}{\beta_b v_a} \cdot \mathsf{C}_{ab\,R}(v_a, \{a_c\}, \{\bar{\kappa}_c\})$$
(2.14)

$$C_{ab\,R}(v_a, \{a_c\}, \{\bar{\kappa}_c\}) = -\int_0^1 \frac{du}{u} \underbrace{\mathbb{R}_{ab}(\{a_c v_a u\}) \mathbb{H}(\{a_c v_a u\}, \{\bar{\kappa}_c\})}_{\text{d.cab.reg}} . \tag{2.15}$$

ccoeff.f90: d_cab_reg

```
! regular contribution for non-zero electron mass
SUBROUTINE c_reg_mass(nni, ia, ib, vp, k2, kb2, betab, mb, cc_r)
       USE physvars
IMPLICIT NONE
          INTEGER,
                                                INTENT(IN)
                                                                :: nni
          INTEGER,
                                                INTENT(IN)
                                                                :: ia
          INTEGER,
                                                INTENT(IN)
          REAL,
                                                INTENT(IN)
                     INTENT(IN)
INTENT(IN)
DIMENSION(1:nni+1), INTENT(IN)
DIMENSION(1:nni+1), INTENT(IN)
          REAL,
          REAL,
REAL,
                                                                   kb2
                                                                :: betab
          REAL,
                                                                :: mb
                                                INTENT (OUT)
          REAL,
         REAL, DIMENSION(1:nni+1) :: ab
INTEGER, PARAMETER :: NR=10 ! integration regions: must be even
INTEGER, PARAMETER :: NR=100 ! integration regions: must be even
REAL, PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad
REAL, PARAMETER :: W13=0.5555555556E0, W2=0.8888888889E0
REAL :: u0, u1, du, u, um, d_cab_reg
ļ
          INTEGER
          ab=SQRT(0.5*betab*mb)*vp/CC
          u0=0.0
!
          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
```

```
cc_r=cc_r+W2*d_cab_reg(u,vp,ia,ib,nni,k2,kb2,betab,mb)
     um=u-du*UPM
     cc_r=cc_r+W13*d_cab_reg(um,vp,ia,ib,nni,k2,kb2,betab,mb)
     um=u+du*UPM
     cc_r=cc_r+W13*d_cab_reg(um, vp, ia, ib, nni, k2, kb2, betab, mb)
  ENDDO
  cc_r=cc_r*du
END SUBROUTINE c_reg_mass
FUNCTION d_cab_reg(u, vp, ia, ib, nni, k2, kb2, betab, mb)
USE mathvars
USE physvars
  IMPLICIT NONE
  REAL,
                                 INTENT(IN)
                                                          [dimensionless]
                                             :: u
                                                        ! Projectile velocity [cm/
                                 INTENT(IN)
  REAL,
                                             :: vp
  INTEGER,
                                 INTENT(IN)
                                                        ! Species number
                                             :: ia
  INTEGER,
                                             :: ib
                                 INTENT(IN)
                                                        ! Species number
  INTEGER,
                                 INTENT(IN)
                                             :: nni
                                                        ! Number of ion species
                                             :: k2
                                                        ! Wave-number squared [1/c
  REAL,
                                 INTENT(IN)
  REAL,
           DIMENSION(1:nni+1), INTENT(IN)
                                             :: kb2
                                                        ! Debye wavenumber squared
  REAL,
           DIMENSION(1:nni+1), INTENT(IN)
                                             :: betab ! Inv temperature array [1
           DIMENSION(1:nni+1), INTENT(IN)
                                                        ! Mass array [keV]
  REAL,
                                             :: mb
  REAL
                                             :: d_cab_reg! [dimensionless]
           DIMENSION(1:nni+1) :: kbar2b, ab, ab2
:: fr, fi, fabs, farg, h, r_ib
  REAL,
  REAL
                                :: kcb, bm_ic, bm_ib, a2_ic, a2_ib, ex, au
  REAL
  INTEGER
  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))
  ! calculate spectral density
  r_{ib}=0
  bm_ib=betab(ib)*mb(ib)
  a2_{ib} = ab(ib)*ab(ib)
  DO ic=1,nni+1
     kcb=kb2(ic)/k2
     bm_ic=betab(ic)*mb(ic)
     a2_ic =ab(ic)*ab(ic)
     IF (ic == ib) THEN
        ex=1.
     ELSE
        au=(a2_ic-a2_ib)*u ! avoids exp of
        ex=EXP(-au)
                             ! large numbers
     ENDIF
     r_ib=r_ib + kcb*SQRT(bm_ic/bm_ib)*ex
  ENDDO
  r_{ib}=1./r_{ib}
  d_cab_reg=-r_ib*h/(u*TWOPI) ! See * in ccoeff_1.0.pdf
END FUNCTION d_cab_reg
```

C. The Singular Contribution: c_sing

The singular contribution,

$$\mathcal{A}_{b,S}^{C} = \left[\frac{e_p^2 \kappa_b^2}{4\pi} \left(\frac{\beta_b m_b}{2\pi} \right)^{1/2} v_p \right] \int_0^1 du \, u^{-1/2} e^{-\frac{1}{2} \beta_b m_b v_p^2 u} \left[-\ln \left\{ \frac{\beta_b e_b e_p}{4\pi} K \frac{m_b}{m_{pb}} \frac{u}{1-u} \right\} - 2\gamma \right],$$
(2.16)

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

$$\int_0^1 du \, u^{-1/2} e^{-\frac{1}{2}\beta_b m_b v_p^2 u} \left[\ln \left\{ \frac{u}{1-u} \right\} - \ln \left\{ \frac{\beta_b e_b e_p}{4\pi} K \frac{m_b}{m_{pb}} \right\} - 2\gamma \right] , \qquad (2.17)$$

which motivates the definition

$$C_{b,S}^{\ell\ell C} = c_{b,1} c_{b,2} \cdot \mathsf{C}_{S}(a_{pb}, b_{pb}) \tag{2.18}$$

$$C_{s}(a,b) = \int_{0}^{1} du \, u^{-1/2} e^{-a \, u} \left[-\ln \left\{ \frac{u}{1-u} \right\} + b \right]$$
 (2.19)

$$a_{pb} = \frac{1}{2} \beta_b m_b v_p^2$$
 and $b_{pb} = -\ln \left\{ \frac{\beta_b e_b e_p}{4\pi} K \frac{m_b}{m_{pb}} \right\} - 2\gamma$ (2.20)

$$c_{b,1} = \frac{e_p^2 \kappa_b^2}{4\pi} \quad c_{b,2} = \left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} v_p .$$
 (2.21)

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

```
acoeff.f90: sing
```

```
! singular contribution for non-zero electron mass
SUBROUTINE c_sing_mass(a, b, cc_s)
                INTENT(IN)
INTENT(IN)
       REAL,
       REAL,
                            :: b
       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
                PARAMETER :: W13=0.555555556E0, W2=0.8888888889E0
             _s=0
          CC.
          u0 = 0
          u1=1
          du=(u1-u0)/NS
          u=u0-du
          DO iu=1,NS,2 ! Gaussian quadrature
             u=u+2.E0*du
             cc_s=cc_s+W2*dcab_sing(u,a,b)
             um=u-du*UPM
             cc_s=cc_s+W13*dcab_sing(um,a,b)
             um=u+du*UPM
             cc_s=cc_s+W13*dcab_sing(um,a,b)
          cc_s=cc_s*du
     END SUBROUTINE c_sing_mass
 a = betab * mb * vp^2/ 2
     FUNCTION dcab_sing(u, a, b)
       IMPLICIT NONE
       REAL,
                    INTENT(IN)
                                               [dimensionless]
                                              [dimensionless]
       REAL,
                    INTENT(IN)
                                            ! a=(1/2)*beta*mpc2*vp^2/C^2
       REAL,
                    INTENT(IN)
                                            ! [dimensionless]
                                :: dcab_sing! [dimensionless]
       R.F.AT.
       dcab_sing=EXP(-a*u)*(-LOG(u/(1-u)) + b)/SQRT(u) ! Eq BPS (9.5)
     END FUNCTION dcab_sing
```

D. The Quantum Correction: c_quantum

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

$$C_b^{\ell\ell \,\text{QM}} = -\frac{e_p^2 \,\kappa_b^2}{4\pi} \, \left(\frac{\beta_b m_b}{2\pi}\right)^{1/2} \int_0^\infty du \, \left[\text{Re} \, \psi \left\{ 1 + i \, \frac{\bar{\eta}_{pb}}{u} \right\} - \ln \left\{ \frac{\bar{\eta}_{pb}}{u} \right\} \right] \left[e^{-\frac{1}{2} \, \beta_b m_b v_p^2 (u-1)^2} - e^{-\frac{1}{2} \, \beta_b m_b v_p^2 (u+1)^2} \right] \,. \tag{2.22}$$

The quantum function we need to code is therefore

$$C_b^{\ell\ell_{\rm QM}} = \underbrace{** \left[\frac{e_p^2 \kappa_b^2}{4\pi} \left(\frac{\beta_b m_b}{2\pi} \right)^{1/2} v_p \right]}_{C_b \ 1 \ C_b \ 2} \cdot \mathsf{C}_1^{\mathrm{QM}}(a_{pb}, \tilde{\eta}_{pb}) , \qquad (2.23)$$

where the arguments of the function are defined by

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

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

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

$$(2.25)$$

and the function itself takes the form

$$\mathsf{C}_{1}^{\mathrm{QM}}(a,\eta) = -\int_{0}^{\infty} du \left[\operatorname{Re} \psi \left\{ 1 + i \frac{\eta}{u} \right\} - \ln \left\{ \frac{\eta}{u} \right\} \right] \frac{1}{2a \, u} \left[e^{-a(u-1)^{2}} - e^{-a(u+1)^{2}} \right] . \tag{2.26}$$

acoeff.f90: quantum

```
quantum contribution for non-zero electron mass
SUBROUTINE c_quantum_mass(ia, ib, a, eta, aq)
IMPLICIT NONE
         INTEGER, INTENT(IN)
                                  :: ia
                                             ! species index
                                  :: ib
                                             ! species index
         INTEGER, INTENT(IN)
                                           ! [dimensionless] (1/2) betab mb vp^2
         REAL,
                    INTENT(IN) :: a
                    INTENT(IN) :: eta
                                            ! [dimensionless] ep eb/4pi hbar vp
         REAL,
         REAL, INTENT(OUT) :: aq
REAL :: u0, u1, du, u, um
INTEGER, PARAMETER :: NQ=1000 ! integration regions quantum : must
REAL, PARAMETER :: UPM=0.7745966692E0 ! parameters for Gaussian Quad
REAL, PARAMETER :: W13=0.5555555556E0, W2=0.8888888889E0
         REAL,
         REAL :: dcq
INTEGER :: iu
         ! choose plot range of gaussian e^{-a^2}
         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
  ! gaussian quadrature
  du=(u1-u0)/NQ
  u=u0-du
  DO iu=1,NQ,2 ! Gaussian quadrature u=u+2.E0*du
      aq=aq+W2*dcq(u,a,eta)
      um=u-du*UPM
      aq=aq+W13*dcq(um,a,eta)
      um=u+du*UPM
      aq=aq+W13*dcq(um,a,eta)
  ENDDÓ
aq=aq*du
END SUBROUTINE c_quantum_mass
FUNCTION dcq(u, a, eta)
USE physvars
IMPLICIT NONE
  REAL,
                                       INTENT(IN)
INTENT(IN)
                                                                           [dimensionless]
                                                      :: u
:: a
  REAL,
  REAL,
                                                                           [dimensionless]
                                       INTENT(IN)
                                                      :: eta
                                                      :: dcq ! [dimensionless]
  REAL
  REAL
                       :: repsi, au, eu, ep, em, psilog, ch, sh, csh
  eu = eta/u

au = 2*a*u
  psilog = repsi(eu) - LOG(eu)

em = EXP(-a * (u - 1)**2)

ep = EXP(-a * (u + 1)**2)
  csh = em - ep
  dcq =-psilog*csh/au
END FUNCTION dcq
```

Appendix A: Calculating the Real and Imaginary Parts of F

We can write the dielectric function (1.6) as a sum over plasma components,

$$F(v) = \sum_{b} F_b(v) , \qquad (A1)$$

where we express the contribution from plasma species b as

$$F_b(v) = -\int_{-\infty}^{\infty} du \, \frac{\rho_b(v)}{v - u + i\eta} \tag{A2}$$

$$\rho_b(v) = \kappa_b^2 \sqrt{\frac{\beta_b m_b}{2\pi}} v \exp\left\{-\frac{1}{2} \beta_b m_b v^2\right\}. \tag{A3}$$

We will often decompose F into its contribution from electrons and ions and write

$$F(v) = F_e(v) + F_{\rm I}(v) . \tag{A4}$$

Note the reflection property

$$F_b(-v) = F_b^*(v) , \qquad (A5)$$

which means that the real part of $F_b(v)$ is even in v and the imaginary part is odd. For numerical work it is best to use the explicit real and imaginary parts of F, which can be written

$$F_{\text{Re}}(v) = \sum_{b} \kappa_b^2 \left[1 - 2\sqrt{\frac{\beta_b m_b}{2}} v \operatorname{daw} \left\{ \sqrt{\frac{\beta_b m_b}{2}} v \right\} \right]$$
(A6)

$$F_{\text{Im}}(v) = \sqrt{\pi} \sum_{b} \kappa_b^2 \sqrt{\frac{\beta_b m_b}{2}} \quad v \exp\left\{-\frac{\beta_b m_b}{2} v^2\right\} = \pi \rho_{\text{tot}}(v) , \qquad (A7)$$

where the Dawson integral is defined by

$$daw(x) = \int_0^x dy \, e^{y^2 - x^2} = \frac{\sqrt{\pi}}{2} \, e^{-x^2} erfi(x) . \tag{A8}$$

The limits of small and large arguments of the Dawson function are

$$daw(x) = x + \frac{2x^3}{3} + \frac{4x^5}{15} + \mathcal{O}(x^7)$$
(A9)

$$daw(x) = \frac{1}{2x} + \frac{1}{4x^3} + \frac{3}{8x^5} + \mathcal{O}(x^{-7}) . \tag{A10}$$

The functions $F_b(v)$ have units of wave-number-squared $[1/L^2]$ and their argument v has units of velocity. We can express the functions $F_b(v)$ in terms of a single dimensionless function $\mathbb{F}(x)$ as follows:

$$F_b(v) = \kappa_b^2 \mathbb{F}\left(\sqrt{\frac{\beta_b m_b}{2}} v\right) \tag{A11}$$

with

$$\mathbb{F}(x) = \int_{-\infty}^{\infty} dy \, \frac{\bar{\rho}(y)}{y - x - i\eta} \tag{A12}$$

$$\bar{\rho}(y) = \frac{1}{\sqrt{\pi}} y e^{-y^2} .$$
 (A13)

Relation (A11) holds because $\rho_b(u) = \kappa_b^2 \bar{\rho}(y)$ for $u = (2/\beta_b m_b)^{1/2} y$. The reflection property (A5) becomes

$$\mathbb{F}(-x) = \mathbb{F}^*(x) , \qquad (A14)$$

which means that the real part is even in x and the imaginary part is odd,

$$\mathbb{F}_{Re}(-x) = \mathbb{F}_{Re}(x) \tag{A15}$$

$$\mathbb{F}_{\text{Im}}(-x) = -\mathbb{F}_{\text{Im}}(x) . \tag{A16}$$

As with expressions (A6) and (A7), the real and imaginary parts can be written

$$\mathbb{F}_{\text{Re}}(x) = 1 - 2x \operatorname{daw}(x) \tag{A17}$$

$$\mathbb{F}_{\text{Im}}(x) = \pi \,\bar{\rho}(x) = \sqrt{\pi} \,x \,e^{-x^2} \,.$$
 (A18)

Let us now establish the forms (A6) and (A7) for the real and imaginary parts of F(v). Staring with

$$\frac{1}{y-x-i\eta} = \mathsf{P}\frac{1}{y-x} + i\pi\,\delta(y-x)\;,\tag{A19}$$

the imaginary part becomes

$$\mathbb{F}_{\text{Im}}(x) = \int_{-\infty}^{\infty} dy \operatorname{Im} \frac{\bar{\rho}(y)}{y - x - i\eta} = \int_{-\infty}^{\infty} dy \, \bar{\rho}(y) \, \pi \delta(y - x) = \pi \bar{\rho}(x) . \tag{A20}$$

The real part of the function must be evaluated by a principal part integral,

$$\mathbb{F}_{\text{Re}}(x) = \mathsf{P} \int_{-\infty}^{\infty} dy \, \frac{\bar{\rho}(y)}{y - x} \,. \tag{A21}$$

Let us add and subtract unity in the form

$$\int_{-\infty}^{\infty} \frac{dy}{y} \,\bar{\rho}(y) = 1 \,\,, \tag{A22}$$

so that

$$\mathbb{F}_{\text{Re}}(x) = 1 + \mathsf{P} \int_{-\infty}^{\infty} dy \left[\frac{\bar{\rho}(y)}{y - x} - \frac{\bar{\rho}(y)}{y} \right] \tag{A23}$$

$$= 1 + \mathsf{P} \int_{-\infty}^{\infty} dy \, \frac{x}{y(y-x)} \, \bar{\rho}(y) \tag{A24}$$

$$= 1 + \frac{x}{\sqrt{\pi}} P \int_{-\infty}^{\infty} dy \, \frac{e^{-y^2}}{y - x} \,. \tag{A25}$$

Making the change of variables y' = y - x (and dropping the prime) we can write

$$\mathbb{F}_{\text{Re}}(x) = 1 + \frac{x}{\sqrt{\pi}} P \int_{-\infty}^{\infty} \frac{dy}{y} e^{-(y+x)^2}$$
(A26)

$$= 1 + \frac{x e^{-x^2}}{\sqrt{\pi}} P \int_{-\infty}^{\infty} \frac{dy}{y} e^{-y^2 - 2xy}$$
 (A27)

$$= 1 + \frac{x e^{-x^2}}{\sqrt{\pi}} \lim_{\epsilon \to 0^+} \left[\int_{-\infty}^{-\epsilon} \frac{dy}{y} e^{-y^2 - 2xy} + \int_{\epsilon}^{\infty} \frac{dy}{y} e^{-y^2 - 2xy} \right] . \tag{A28}$$

In the last expression we have used the definition of the principal part integration. Making a change of variables y' = -y in the first integral in square brackets gives (and again dropping the prime)

$$\int_{-\infty}^{-\epsilon} \frac{dy}{y} e^{-y^2 - 2xy} = -\int_{\epsilon}^{\infty} \frac{dy}{y} e^{-y^2 + 2xy} , \qquad (A29)$$

and this allows us to write

$$\mathbb{F}_{\text{Re}}(x) = 1 - \frac{x e^{-x^2}}{\sqrt{\pi}} \lim_{\epsilon \to 0^+} \int_{\epsilon}^{\infty} \frac{dy}{y} e^{-y^2} \left[e^{2xy} - e^{-2xy} \right]. \tag{A30}$$

The term in square braces is just $2\sinh(2xy)$, which renders the factor 1/y harmless when the limit $\epsilon \to 0^+$ is taken,

$$\mathbb{F}_{\text{Re}}(x) = 1 - \frac{2x e^{-x^2}}{\sqrt{\pi}} \int_0^\infty dy \, e^{-y^2} \, \frac{\sinh 2xy}{y} = 1 - 2x \, \text{daw}(x) \,. \tag{A31}$$

The latter form hold because this is just another integral representation of the Dawson function,

$$daw(x) = \frac{e^{-x^2}}{\sqrt{\pi}} \int_0^\infty dy \, e^{-y^2} \, \frac{\sinh 2xy}{y} \, . \tag{A32}$$

Compare this with

$$daw(x) = e^{-x^2} \int_0^x dy \, e^{y^2} \,. \tag{A33}$$