

# P2Scattering.m v1.3

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ABSTRACT: The MATHEMATICA package `P2Scattering.m`, first released alongside our work with Pierrick Bousseau and Pierre Descombes [1], provides a suite of routines to study the scattering diagram for the derived category  $\mathcal{C} = D^b(K_{\mathbb{P}^2})$  of coherent sheaves on local  $\mathbb{P}^2$  along the large volume, orbifold and  $\Pi$ -stability slices in the space of Bridgeland stability conditions, and extracting the corresponding generalized Donaldson-Thomas invariants  $\Omega_Z(\gamma)$ . We provide a list of all routines and give a few examples.

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## 1. Summary

The MATHEMATICA package `P2Scattering.m` provides a suite of routines to study the scattering diagram for the derived category  $\mathcal{C} = D^b(K_{\mathbb{P}^2})$  of coherent sheaves on local  $\mathbb{P}^2$  along the large volume, orbifold and  $\Pi$ -stability slices in the space of Bridgeland stability conditions, and extracting the corresponding generalized Donaldson-Thomas invariants  $\Omega_Z(\gamma)$ .

Following [?], the charge vector  $\gamma$  stands either for the Chern vector  $[r, d, \text{ch}_2]$ , its integral cousin  $[r, d, \chi]$  where  $\chi = r + \frac{3}{2}d + \text{ch}_2$ , or the dimension vector  $(n_1, n_2, n_3)$  associated to the tilting sequence

$$(E_1 = i_*(\mathcal{O}), \quad E_2 = i_*(\Omega(1))[1], \quad E_3 = i_*(\mathcal{O}(-1))[2]) \quad (1.1)$$

such that

$$(n_1, n_2, n_3) = \left(-\frac{3}{2}d - \text{ch}_2 - r, -\frac{1}{2}d - \text{ch}_2, \frac{1}{2}d - \text{ch}_2\right) = (-\chi, r + d - \chi, r + 2d - \chi) \quad (1.2)$$

The central charge is given by  $Z(\gamma) = -rT + dT_D - \text{ch}_2$ , where  $(T, T_D)$  are

$$T = s + it, \quad T_D = \frac{1}{2}(s + it)^2 \quad (1.3)$$

for the large volume slice, or

$$\begin{pmatrix} T \\ T_D \end{pmatrix} = \begin{pmatrix} -\frac{1}{2} \\ \frac{1}{3} \end{pmatrix} + \int_{\tau_o}^{\tau} \begin{pmatrix} 1 \\ u \end{pmatrix} C(u) du \quad (1.4)$$

for the  $\Pi$ -stability slice, where  $C(\tau) = \frac{\eta(\tau)^9}{\eta(3\tau)^3}$  is a weight 3 Eisenstein series for  $\Gamma_1(3)$  and  $\tau_o = -\frac{1}{2} + \frac{i}{2\sqrt{3}}$  is the orbifold point. The geometric rays in the scattering diagram are then the real codimension-one loci

$$\mathcal{R}_\psi(\gamma) = \{\tau : \Re(e^{i\psi} Z(\gamma)) = 0\} \quad (1.5)$$

For the large volume central charge and  $\psi = 0$ , the scattering diagram reduces to the one constructed in [1]. Finally, around the orbifold slice, we consider a two-dimensional slice  $\theta_1 + \theta_2 + \theta_3 = -\sin \epsilon$  in the space of King stability parameters  $\theta_i = \Im(e^{-i\epsilon} Z_\tau(\text{ch } E_i))$ , and the rays are the real-codimension one loci  $n_1\theta_1 + n_2\theta_2 + n_3\theta_3 = 0$ .

The package file `P2Scattering.m` and various example files can be obtained from <https://github.com/bpioline/P2Scattering>

## 1.1 Basic usage

Assuming that the file `P2Scattering.m` is present in the user's MATHEMATICA Application directory, the package is loaded by entering

```
In[1]:= <<P2Scattering'
Out[1]:= P2Scattering 1.2 -- A package for evaluating DT invariants
on  $K_{P^2}$ 
```

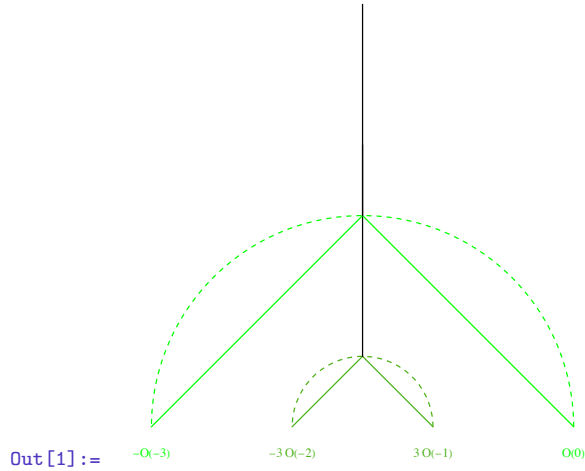
If the file `P2Scattering.m` has not yet been copied in the user's MATHEMATICA Application directory but is in the same directory as the notebook, evaluate instead

```
In[1]:= SetDirectory[NotebookDirectory[]]; <<P2Scattering'
Out[1]:= P2Scattering 1.2- A package for evaluating evaluating DT
invariants on  $K_{P^2}$ 
```

For given charge  $\gamma = [r, d, \chi]$  and point  $(s, t)$  on the large volume slice, the trees contributing to the index  $\Omega_{(s,t)}(\gamma)$  can be found by using the routine `ScanAllTrees`, for example for  $\gamma = [3, 0, 0]$  through the point  $(s, t) = (-\frac{3}{2}, 2)$ ,

```
In[1]:= LiTrees=ScanAllTrees[{0, 3, 0}, {-3/2, 2}]
Out[1]:= {{-3 Ch[-2], 3 Ch[-1]}, {-Ch[-3], Ch[0]}}
```

```
In[1]:= ScattDiagLV[LiTrees, 0]
```



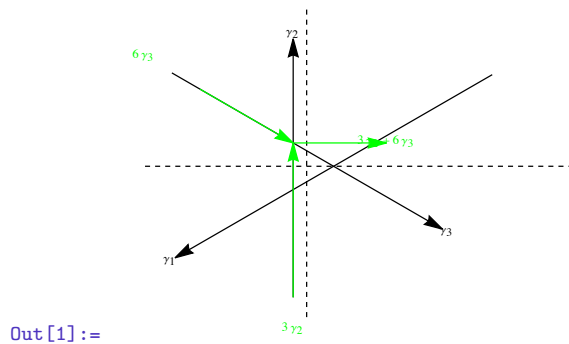
```
In[1]:= Limit[EvaluateKronecker[ScattIndex[LiTrees]], y -> 1]
Out[1]:= {18, 9}
```

reproducing the GV invariant  $N_3^0 = 27$ . Similarly, one can find the trees contributing near the orbifold point using `McKayScanAllTrees`: for the same charge, corresponding to dimension vector  $(0, 3, 6)$ , a single tree contributes in the anti-attractor chamber, with index 18,

```
In[1]:= LiTrees = McKayScanAllTrees[chiton[{0, 3, 0}]]; LiTrees /.
McKayrep
Out[1]:= {{3γ2, 6γ3}}
```

```
In[1]:= Limit[EvaluateKronecker[McKayScattIndex[LiTrees]], y -> 1]
Out[1]:= {18}
```

```
In[1]:= Show[McKayInitialRays[2], McKayScattDiag[LiTrees]]
```



More examples can be found on the GitHub repository.

## 1.2 Online documentation

The package contains many more routines, documented below, which can be used independently. Basic inline documentation can be obtained by typing e.g.

```
In[1]:= ?EichlerT
Out[1]:= EichlerT[tau_] gives numerical value of T[tau] using Eichler
integral, by mapping back to fundamental domain  $F_C$ 
```

## 1.3 History

The first version of this package was released together with the preprint [?].

## 2. Symbols and global variables

- **tau**: Kähler modulus in Poincaré upper half-plane
- **tau1**: Real part of **tau**
- **tau2**: Imaginary part of **tau**
- **taup**: Fricke transform  $\tau' = -1/(3\tau)$
- **y**: refinement parameter
- **a**: Parameter running from 0 to 1 along rays
- **Ch[m\_]**: denotes the charge vector for  $\mathcal{O}(m)$  if  $m$  is integer; More generally, if  $m$  is rational, it denotes the charge of the image of  $\mathcal{O}$  which is massless at  $\tau = m$
- **Ch[m\_][n\_]**: denotes the charge vector for  $\mathcal{O}(m)[n]$
- **Kr\_m[p\_,q\_]**: denotes the index of the Kronecker quiver with  $m$  arrows, dimension vector  $(p, q)$
- **gam1**: Charge vector  $[-1, 0, -1]$  for  $E_1[-1] = \mathcal{O}(0)[-1]$
- **gam2**: Charge vector  $[2, -1, 0]$  for  $E_2[-1] = \Omega(1)[0]$
- **gam3**: Charge vector  $[-1, 1, 0]$  for  $E_3[-1] = \mathcal{O}(-1)[1]$
- **tau0**: orbifold point  $-1/2 + i/(2\sqrt{3})$
- **QuantumVolume**: Quantum period  $\mathcal{V} = \Im T(0) \simeq 0.462758$
- **MLV**: Monodromy matrix in  $[r, d, \chi]$  basis around  $\tau = i\infty$
- **MCon**: Monodromy matrix in  $[r, d, \chi]$  basis around at  $\tau = 0$
- **MOrb**: Monodromy matrix in  $[r, d, \chi]$  basis around orbifold point  $\tau_o$
- **MOrbp**: Monodromy matrix in  $[r, d, \chi]$  basis around orbifold point  $\tau_o + 1$
- **Trees[r\_,d\_,chi\_]**: gives the list of precomputed trees at large volume, when available
- **McKayTrees[{n1\_,n2\_,n3\_}]**: gives the list of precomputed trees around the orbifold point, when available
- **ExcepSlopes**: List of slopes of exceptional bundles between -3 and 4
- **ListSubsetsAny**: Precomputed list of all binary splits of  $\text{Range}[n]$  for  $n=2..10$ , used by **ListStableTrees**
- **FourierC**: List of the first 50 Fourier coefficients of Eisenstein series  $C(\tau)$
- **FourierCp**: List of the first 50 Fourier coefficients of Eisenstein series  $C'(\tau')$

### 3. Operations on Chern vectors and dimension vectors

- `Euler`[ $\{r\_ , d\_ , chi\_ \}, \{rr\_ , dd\_ , cchi\_ \}$ ]: computes the Euler form on  $D^b(\mathbb{P}^2)$
- `DSZ`[ $\{r\_ , d\_ , chi\_ \}, \{rr\_ , dd\_ , cchi\_ \}$ ]: computes the antisymmetrized Euler form  $3(rd' - r'd)$  on  $D^b(\mathbb{P}^2)$
- `McKayDSZ`[ $\{n1\_ , n2\_ , n3\_ \}, \{nn1\_ , nn2\_ , nn3\_ \}$ ]: computes the antisymmetrized quiver Euler form
- `Disc`[ $\{r\_ , d\_ , chi\_ \}$ ]: computes the discriminant  $\Delta(\gamma) = d^2 - 2r(\chi - r - \frac{3}{2}d)/(2r^2)$
- `Disch2`[ $\{r\_ , d\_ , ch2\_ \}$ ]: computes the discriminant  $\Delta(\gamma) = (d^2 - 2r \text{ch}_2)/(2r^2)$
- `DiscR`[ $\{r\_ , d\_ , chi\_ \}$ ]: computes the rescaled discriminant  $r^2\Delta(\gamma) = d^2 - 2r(\chi - r - \frac{3}{2}d)$
- `GenSlope`[ $\{r\_ , d\_ , chi\_ \}$ ]: computes the slope  $d/r$  if  $r \neq 0$ , or  $\chi/d - \frac{3}{2}$  if  $r = 0$
- `DimGieseker`[ $\{r\_ , d\_ , chi\_ \}$ ]: computes expected dimension of moduli space of Gieseker-semistable sheaves
- `DimMcKay`[ $\{n1\_ , n2\_ , n3\_ \}$ ]: computes the dimension of quiver moduli space in anti-attractor chamber
- `ch2tochi`[ $r\_ , d\_ , ch2\_$ ]: produces corresponding  $[r, d, \chi]$
- `chitoch2`[ $r\_ , d\_ , chi\_$ ]: produces corresponding  $[r, d, \text{ch}_2]$
- `chiton`[ $\{r\_ , d\_ , chi\_ \}$ ]: produces the corresponding dimension vector  $(n_1, n_2, n_3)$
- `ntochi`[ $\{n1\_ , n2\_ , n3\_ \}$ ]: produces the corresponding charge vector  $[r, d, \text{chi}]$
- `SpecFlow`[ $\{r\_ , d\_ , ch2\_ \}, k\_$ ]: computes the translated charge vector  $[r, d, \chi](k)$
- `SpecFlowch2`[ $\{r\_ , d\_ , ch2\_ \}, k\_$ ]: computes the translated charge vector  $[r, d, \text{ch}_2](k)$
- `repCh`[: Replacement rule mapping  $\text{Ch}[s]$  or  $\text{Ch}[s][n]$  to their charge vector  $[r, d, \chi]$
- `repCh2`[: Replacement rule mapping  $\text{Ch}[s]$  or  $\text{Ch}[s][n]$  to their charge vector  $[r, d, \text{ch}_2]$
- `repChn`[: Replacement rule mapping  $Ch[m]$  into `chiton`[ $\{1, m, 1+m(m+3)/2\}$ ]
- `repCh0`[: Replacement rule mapping  $Ch[m]$  to string  $O(m)$
- `repKr`[: replaces  $Kr_m[p, q]$  by 1
- `LPCurve`[ $\mu\_$ ]: computes the Drezet-Le Potier curve  $\delta(\mu)$
- `ExceptToChi`[ $\mu\_$ ]: gives the Chern vector  $[r, d, \chi]$  of the exceptional bundle of slope  $\mu$
- `CPointchi`[ $\tau\_$ ]: gives the charge vector  $[r, d, \chi]$  of an object that becomes massless at  $\tau$  (assuming  $\tau$  is a rational number)
- `CPointch2`[ $\tau\_$ ]: gives the charge vector  $[r, d, \text{ch}_2]$  of an object that becomes massless at  $\tau$  (assuming  $\tau$  is a rational number)
- `EvaluateKronecker`[ $f\_$ ]: replaces each  $Kr_m[p, q]$  with the index of the Kronecker quiver with  $m$  arrows and dimension vector  $(p, q)$ , using routines taken from `CoulombHiggs.m` package
- `McKayrep`[: replaces  $\{n1\_ , n2\_ , n3\_ \}$  by  $n_1\gamma_1 + n_2\gamma_1 + n_3\gamma_3$

#### 4. Kähler moduli space and central charge for $\Pi$ -stability

- `ToFundDomain0[tau_]`: produces  $\{\tau', M'\}$  such that  $M' \cdot \tau' = \tau$  and  $\tau'$  lies in fundamental domain of  $\Gamma_1(3)$  centered around orbifold
- `ToFundDomainC[tau_]`: produces  $\{\tau', M'\}$  such that  $M' \cdot \tau' = \tau$  and  $\tau'$  lies in fundamental domain of  $\Gamma_1(3)$  centered around conifold
- `ToFundDomainCSeq[tau_]`: produces  $\{\tau', M'\}$  such that  $M' \cdot \tau' = \tau$  and  $\tau'$  lies in fundamental domain of  $\Gamma_1(3)$  centered around conifold,  $M$  is a string of  $U, V, T[m]$  generators
- `ToFundDomain0Approx[tau_, precision_]`: produces  $\{\tau', M'\}$  such that  $M' \cdot \tau' = \tau$  and  $\tau'$  lies in fundamental domain of  $\Gamma_1(3)$  centered around orbifold
- `ToFundDomainCApprox[tau_, precision_]`: produces  $\{\tau', M'\}$  such that  $M' \cdot \tau' = \tau$  and  $\tau'$  lies in fundamental domain of  $\Gamma_1(3)$  centered around conifold
- `ApplyGamma13Lift[M_, tau_]`: produces the image of  $\tau$  under the 3x3 monodromy matrix  $M$
- `MonodromyOnCharge[M_, {r_, d_, chi_}]`: computes the image of charge vector  $[r, d, \chi]$  under sequence of monodromies
- `MonodromyOnTau[M_, tau_]`: computes the image of  $\tau$  under sequence of monodromies
- `FundDomain0[k_]`: produces the Graphics directives for the fundamental domain of  $\Gamma_1(3)$  on the interval  $[-\frac{1}{2} + k, \frac{1}{2} + k]$
- `FundDomainC[k_]`: produces the Graphics directives for the fundamental domain of  $\Gamma_1(3)$  on the interval  $[k, k + 1]$
- `FundDomain3[k_]`: produces the Graphics directives for the fundamental domain of  $\Gamma_1(3) + \mathbb{Z}_3$  images on the interval  $[k, k + 1]$
- `FundDomainRulesC[k_]`: gives a list of rules  $\tau \rightarrow \tau(a)$  for boundaries of `FundDomainC[k]` parametrized by  $0 < a < 1$
- `FundDomainRules0[k_]`: gives a list of rules  $\tau \rightarrow \tau(a)$  for boundaries of `FundDomain0[k]` parametrized by  $0 < a < 1$
- `FundDomainRules3[k_]`: gives a list of rules  $\tau \rightarrow \tau(a)$  for boundaries of `FundDomain3[k]` parametrized by  $0 < a < 1$
- `EichlerC[tau_]`: numerically evaluates the Eisenstein series  $C(\tau)$
- `EichlerCp[taup_]`: numerically evaluates the Eisenstein series  $C'(\tau')$
- `Eichlerf1[tau_]`: evaluates  $2\pi i(\tau + 1/2) + \bar{f}_1(\tau)$
- `Eichlerf2[tau_]`: evaluates  $\frac{1}{2}(2\pi i)^2(\tau + \frac{1}{2})^2 + 2\pi i(\tau + \frac{1}{2})\bar{f}_1(\tau) + \bar{f}_2(\tau)$
- `Eichlerf1b[tau_]`:  $= \sum_{n \geq 1} \frac{c_n}{n} e^{2\pi i n \tau}$

- `Eichlerf2b[tau_]`:  $:= -\sum_{n \geq 1} \frac{c_n}{n^2} e^{2\pi i n \tau}$
- `Eichlerf1p[taup_]`:  $:= \sum_{n \geq 1} \frac{c_n}{n} e^{2\pi i n \tau}$
- `Eichlerf2p[taup_]`:  $:= 2\pi i \tau' f_1'(\tau') - \sum_{n \geq 1} \frac{c_n}{n^2} e^{2\pi i n \tau}$
- `EichlerTLV[tau_]`: gives numerical value of  $T(\tau)$  using Eichler integral near LV point
- `EichlerTDLV[tau_]`: gives numerical value of  $T_D(\tau)$  using Eichler integral near LV point
- `EichlerTC[tau_]`: gives numerical value of  $T(\tau)$  using Eichler integral near conifold point
- `EichlerTDC[tau_]`: gives numerical value of  $T_D(\tau)$  using Eichler integral near conifold point
- `EichlerZ[{r_,d_,chi_},tau_]`: gives numerical value of  $Z_\tau(\gamma)$  by mapping  $\tau$  back to fundamental domain  $\mathcal{F}_C$
- `EichlerZch2[{r_,d_,ch2_},tau_]`: gives numerical value of  $Z_\tau(\gamma)$  by mapping  $\tau$  back to fundamental domain  $\mathcal{F}_C$
- `EichlerZch2LV[{r_,d_,ch2_},tau_]`: gives numerical value of  $Z_\tau(\gamma)$  using Eichler integral at large volume
- `EichlerT[tau_]`: gives numerical value of  $T(\tau)$  using Eichler integral, by mapping back to fundamental domain  $\mathcal{F}_C$
- `EichlerTD[tau_]`: gives numerical value of  $T_D(\tau)$  using Eichler integral, by mapping back to fundamental domain  $\mathcal{F}_C$
- `EichlerTtilde[tau_]`: gives numerical value of  $\tilde{T} = \frac{1}{2\sqrt{3}}(T - \frac{1}{2})$  using Eichler integral, by mapping back to fundamental domain  $\mathcal{F}_C$
- `EichlerTDtilde[tau_]`: gives numerical value of  $\tilde{T}_D = T_D - \frac{1}{2}T - \frac{1}{12}$  using Eichler integral, by mapping back to fundamental domain  $\mathcal{F}_C$



## 5. Large volume scattering diagram

- **ZLV**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, \{s<sub>-</sub>, t<sub>-</sub>\}]: computes the large volume central charge  $-\frac{1}{2}r(s+it)^2 + d(s+it) - (r - \frac{3}{2}d - \chi)$
- **ZLVch2**[\{r<sub>-</sub>, d<sub>-</sub>, ch2<sub>-</sub>\}, \{s<sub>-</sub>, t<sub>-</sub>\}]: computes the central charge  $-\frac{1}{2}r(s+it)^2 + d(s+it) - \text{ch}_2$
- **Wall**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, \{rr<sub>-</sub>, dd<sub>-</sub>, cchi<sub>-</sub>\}, \{s<sub>-</sub>, t<sub>-</sub>\}]: computes  $\Im[Z(\gamma)\overline{Z(\gamma')}]$
- **Wallch2**[\{r<sub>-</sub>, d<sub>-</sub>, ch2<sub>-</sub>\}, \{rr<sub>-</sub>, dd<sub>-</sub>, cch2<sub>-</sub>\}, \{s<sub>-</sub>, t<sub>-</sub>\}]: computes  $\Im[Z(\gamma)\overline{Z(\gamma')}]$
- **TreeCharge**[tree<sub>-</sub>]: computes the total charge  $[r, d, \chi)$  carried by tree (or list of trees).
- **TreeChargech2**[tree<sub>-</sub>]: computes the total charge  $[r, d, \text{ch}_2]$  carried by tree (or list of trees).
- **TreeTop**[tree<sub>-</sub>]: computes the (s,t) coordinate of the root of the tree
- **TreeConstituents**[tree<sub>-</sub>]: gives the flattened list of constituents of Tree
- **FlipTree**[tree<sub>-</sub>]: constructs the reflected tree under  $Ch[m] \rightarrow -Ch[-m]$
- **ShiftTree**[tree<sub>-</sub>, k<sub>-</sub>]: constructs the shifted tree under  $Ch[m] \rightarrow Ch - m + k$
- **ScattCheck**[tree<sub>-</sub>]: returns  $\{\text{charge}, \{x, y\}\}$  of the root vertex if **Tree** is consistent, otherwise  $\{\text{totalcharge}, \{\}\}$
- **ScattSort**[Litree<sub>-</sub>]: sorts trees by growing radius of wall
- **ScattGraph**[tree<sub>-</sub>]: extracts the list of vertices and adjacency matrix of Tree
- **xytost**[\{x<sub>-</sub>, y<sub>-</sub>\}]: computes  $\{x, \sqrt{x^2 + 2y}\}$
- **sttoxy**[\{s<sub>-</sub>, t<sub>-</sub>\}]: computes  $\{s, -\frac{1}{2}(s^2 - t^2)\}$
- **IntersectRays**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, \{rr<sub>-</sub>, dd<sub>-</sub>, cchi<sub>-</sub>\}, z<sub>-</sub>, zz<sub>-</sub>]: returns intersection point (x, y) of two rays if the intersection point lies upward from z and z', or the empty set otherwise; the arguments z, z' can be omitted.
- **IntersectRaysSt**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, \{rr<sub>-</sub>, dd<sub>-</sub>, cchi<sub>-</sub>\}, psi<sub>-</sub>]: returns intersection point (s, t) for scattering rays with phase  $\psi$ , or the empty set if they are collinear
- **TestBranch**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, s<sub>-</sub>]: returns **True** if (s, ·) is on the branch with  $\Im Z(\gamma) > 0$ , **False** otherwise
- **Rayt**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, s<sub>-</sub>, psi<sub>-</sub>]: computes  $t(s)$  for the ray  $\Re[e^{-i\psi}Z^{\text{LV}}(\gamma)] = 0$
- **Rays**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, t<sub>-</sub>, psi<sub>-</sub>]: computes  $s(t)$  for the ray  $\Re[e^{-i\psi}Z^{\text{LV}}(\gamma)] = 0$
- **Raytch2**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, s<sub>-</sub>, psi<sub>-</sub>]: computes  $t(s)$  for the ray  $\Re[e^{-i\psi}Z^{\text{LV}}(\gamma)] = 0$
- **Raysch2**[\{r<sub>-</sub>, d<sub>-</sub>, chi<sub>-</sub>\}, t<sub>-</sub>, psi<sub>-</sub>]: computes  $s(t)$  for the ray  $\Re[e^{-i\psi}Z^{\text{LV}}(\gamma)] = 0$

- **ListStableTrees**[*LiCh*\_, {*s0*\_, *t0*\_} ]: constructs consistent trees from constituents in *LiCh* of the form  $k_i Ch[m_i]$ , which are stable at  $(s_0, t_0)$
- **ListStableTreesPerturb**[*LiCh*\_, {*s0*\_, *t0*\_} ]: constructs consistent trees from constituents in *LiCh* of the form  $k_i Ch[m_i]$ , which are stable at  $(s_0, t_0)$  after perturbing the  $m_i$ 's
- **ListStablePlanarTrees**[*LiCh*\_, {*s0*\_, *t0*\_} ]: constructs consistent planar trees from constituents in *LiCh* of the form  $k_i Ch[m_i]$ , which are stable at  $(s_0, t_0)$
- **ScanConstituents**[{*r*\_, *d*\_, *chi*\_}, {*mmin*\_, *mmax*\_}, {*nmin*\_, *nmax*\_}, *phimax*\_ ]: searches possible list of constituents with slope in  $[mmin, mmax]$ , number of constituents in  $[nmin, nmax]$ , electric potential less than *phimax* and charges adding up to  $[r, d, \chi)$
- **ScanAllTrees**[{*r*\_, *d*\_, *chi*\_}, {*s*\_, *t*\_} ]: constructs all possible trees with charges adding up to  $[r, d, \chi)$  leading to an outgoing ray through the point  $(s, t)$ ; uses **ScanConstituents** with the most conservative values  $mmin = s - t$ ,  $mmax = s + t$ ,  $nmin = 1$ ,  $nmax = 2\varphi_s(\gamma)$  and  $phimax = \varphi_s(\gamma)$
- **ScanBinarySplits**[{*r*\_, *d*\_, *chi*\_}, {*s0*\_, *t0*\_} ]: produces list of  $[r', d', \chi')$  such that  $[r, d, \chi)$  can split into  $[r', d', \chi')$  along the ray starting at  $(s_0, t_0)$
- **ScanKroneckerSplits**[{*r*\_, *d*\_, *chi*\_} ]: scans all possible pairs  $\{-k' Ch(m'), k Ch(m)\}$  such that each pair scatters into  $[r, d, \chi)$
- **ScattIndex**[*TreeList*\_]: computes the index for each tree in *TreeList*; do not trust the result if internal lines have non-primitive charges !
- **ScattIndexInternal**[*TreeList*\_]: computes total charge, list of Kronecker indices associated to each vertex in *Tree*
- **ScattDiag**[*TreeList*\_]: draws scattering diagram in  $(x, y)$  plane for each tree in *TreeList*
- **ScattDiagOverlay**[*TreeList*\_]: overlays scattering diagrams in  $(x, y)$  plane for each tree in *TreeList*
- **ScattDiagxy**[*TreeList*\_, *psi*\_]: draws scattering diagrams in  $(x, y)$  plane overlaying each tree in *TreeList*, marking the initial points
- **ScattDiagInternal**[*Tree*\_]: constructs a list {total charge, coordinate of root, list of line segments, {min(x), max(x)}} for large volume scattering in (x,y) coordinates; used by **ScattDiag**
- **ScattDiagSt**[*TreeList*\_, *psi*\_]: draws scattering diagram in  $(s, t)$  plane for each trees in *TreeList*, approximating each edge by a straight line and overlaying the various trees; if  $\psi$  is omitted, it is assumed to vanish.
- **ScattDiagInternalSt**[*Tree*\_, *psi*\_]: constructs a list {total charge, coordinate of root, list of line segments, {min(s), max(s)}} for large volume scattering in (s,t) coordinates, , approximating each edge by a straight line; if  $\psi$  is omitted, it is assumed to vanish; used by **ScattDiagSt**

- `ScattDiagLV[TreeList_,psi_]`: draws scattering diagram in  $(s, t)$  in  $(s, t)$  plane for each trees in `TreeList`, using exact hyperbolae for rays and overlaying the various trees
- `ScattDiagInternalLV[Tree_,psi_,Style_]`: constructs total charge, coordinate of root and list of line segments in  $(s, t)$  coordinates,  $\{\min(x), \max(x)\}$ , using `PlotStyle`→`Style` for plotting rays ; used by `ScattDiagLV`
- `ScattDiagLZ[TreeList_]`: draws scattering diagram in Li-Zhao  $(s, q)$  plane for each trees in `TreeList`
- `ScattPhi[TreeList_]`: overlays scattering diagrams in  $(s, \varphi)$  plane for each tree in `TreeList`
- `ScattPhiInternal[Tree_]`: constructs a list  $\{\text{total charge, coordinate of root, list of line segments, } \{\min(s), \max(s)\}\}$  in  $(s, \varphi)$  coordinates; used by `ScattPhi`
- `PlotWallRay[{r_,d_,chi_},{rr_,dd_,cchi_}, psi_,{L1_,L2_,H_}]`: plots the local scattering  $[r, d, \chi] \rightarrow [r', d', \chi'] + [r - r', d - d', \chi - \chi']$  for phase  $\psi$  in range  $L_1 < s < L_2, 0 < t < H$
- `WallCircle[{r_,d_,chi_},{rr_,dd_,cchi_}]`: constructs the graphics directive for the wall of marginal stability in  $(s, t)$  coordinates
- `WallLine[{r_,d_,chi_},{rr_,dd_,cchi_}]`: constructs the graphics directive for the wall of marginal stability in  $(s, q)$  coordinates
- `TreeHue[i_,n_]`: specifies the color for the  $i$ -th tree among a list of  $n$  - can be modified at will
- `InitialLabelPosition[m_]`: returns a position  $(s, t)$  for the label for an initial ray with slope  $m$ ; this position is lowered vertically on each call, using variables `LiSlopes` and `LiHeights` to keep track of earlier calls

## 6. Orbifold scattering diagram

- `McKayRayEq[{n1_,n2_,n3_},{u_,v_}]`: gives the linear form vanishing on the scattering ray
- `McKayVec[McKayVec[{n1_,n2_,n3_}]`: computes the positive vector along the ray
- `McKayRay[{n1_,n2_,n3_},{u_,v_},{vardefk1,k2_},tx_]`: produces an arrow from  $(u, v) + k_1 w$  to  $(u, v) + k_2 w$ , where  $w$  is the positive vector along the ray, decorated with text `tx` at the target
- `McKayScattIndex[TreeList_]`: computes the index for each tree in `TreeList`; do not trust the result if internal lines have non-primitive charges !

- `McKayScattIndexInternal[Tree_]`: computes total charge, list of Kronecker indices associated to each vertex in Tree
- `McKayListAllConsistentTrees[{n1_,n2_,n3_}]`: generates consistent scattering trees with leaves carrying charge  $\{p, 0, 0\}, \{0, p, 0\}, \{0, 0, p\}$  adding up to  $(n_1, n_2, n_3)$ , with non-zero DSZ pairing at each vertex, with distinct support
- `McKayListAllTrees[{n1_,n2_,n3_}]`: generates all trees with leaves carrying charge  $\{p, 0, 0\}, \{0, p, 0\}, \{0, 0, p\}$  adding up to  $(n_1, n_2, n_3)$  and with non-zero DSZ pairing at each vertex
- `McKayScattCheck[Tree_]`: returns  $\{charge, \{u, v\}\}$  of the root vertex if Tree is consistent, otherwise  $\{totalcharge, \{\}\}$
- `McKayScattGraph[Tree_]`: extracts the list of vertices and adjacency matrix of Tree
- `McKayScattDiag[TreeList_]`: draws McKay scattering diagram in  $(u, v)$  plane for each tree in TreeList
- `McKayScattDiagInternal[Tree_]`: constructs total charge, coordinate of root and list of line segments in  $(u, v)$  coordinates; used by `McKayScattDiag`
- `McKayIntersectRays[{n1_,n2_,n3_},{nn1_,nn2_,nn3_}]`: returns the intersection point  $(u, v)$  of two rays, or empty set if they are collinear
- `McKayIntersectRays[{n1_,n2_,n3_},{nn1_,nn2_,nn3_},z_,zz_]`: returns intersection point  $(u, v)$  of two rays if the intersection point lies upward from  $z$  and  $z'$ , or empty set otherwise
- `McKayInitialRays[psi_,L_]`: draws the initial rays in  $(u', v')$  plane, rescaling each arrow by a factor of  $L$ . If the argument  $\psi$  is omitted, it is assumed to be  $\frac{\pi}{2}$ .

## 7. Exact scattering diagram

- `CriticalPsi[mu_]`:  $:= \arctan(\mu/\mathcal{V})$
- `IntersectExactRaysLV[{r_,d_,chi_},{rr_,dd_,cchi_},psi_]`: returns value of  $\tau$  at intersection point of two exact rays using `EichlerTLV` to evaluate the periods, or 0 if they are collinear
- `IntersectExactRaysC[{r_,d_,chi_},{rr_,dd_,cchi_},psi_]`: returns value of  $\tau$  at intersection point of two exact rays using `EichlerTC` to evaluate the periods, or 0 if they are collinear
- `XY[tau_,psi_]`: computes the affine coordinates  $(x, y)$  such that scattering rays are straight lines  $ry + dx - ch_2 = 0$

- **CPointxy**[*tau\_*]: computes the  $(x, y)$  coordinate of initial point  $Ch[\tau]$  (assuming  $\tau$  is a rational number)
- **IntegralCurve**[*tauinit\_*, *tangent\_*, {*ainit\_*, *amin\_*, *amax\_*}, *boundaries\_*]: produces a function  $f : a \in [0, 1] \rightarrow \mathbb{H}$  with  $f(ainit) = tauinit$  following the tangent direction (an expression in  $\tau$ ) and stopping at the boundaries (by default:  $\{Im\tau = 0.01\}$ ). The range of integration parameters  $\{amin, amax\}$  can be infinite provided the actual rays remain finite by hitting the boundaries.
- **NormalizeFunctionDomain**[*fun\_*]: rescales the argument of the **InterpolatingFunction** *fun* to interval  $[0, 1]$
- **DtauT**[*tau\_*]: numerically evaluates  $\partial_\tau T(\tau)$
- **DtauZch2**[{*r\_*, *d\_*, *ch2\_*} *mtau\_*]: numerically evaluates  $\partial_\tau Z_\tau(\gamma)$
- **DtauZ**[{*r\_*, *d\_*, *chi\_*} *tau\_*]: numerically evaluates  $\partial_\tau Z_\tau(\gamma)$
- **ArgDtauT**[*vardeftau*]: computes the argument of  $T'(\tau)$ , between  $-\pi$  and  $\pi$
- **ArgDtauTD**[*vardeftau*]: computes the argument of  $T'_D(\tau)$ , between  $-\pi$  and  $\pi$
- **ArgDtauZch2**[{*r\_*, *d\_*, *ch2\_*} *tau\_*]: computes the argument of  $\partial_\tau Z_\tau(\gamma)$ , between  $-\pi$  and  $\pi$
- **UnitDtauT**[*vardeftau*]: numerically evaluates  $\partial_\tau T(\tau)/|\partial_\tau T(\tau)|$
- **UnitDtauTD**[*vardeftau*]: numerically evaluates  $\partial_\tau T_D(\tau)/|\partial_\tau T_D(\tau)|$
- **UnitDtauZ**[{*r\_*, *d\_*, *chi\_*} *tau\_*]: numerically evaluates  $\partial_\tau Z_\tau(\gamma)/|\partial_\tau Z_\tau(\gamma)|$
- **UnitDtauZch2**[{*r\_*, *d\_*, *ch2\_*} *tau\_*]: numerically evaluates  $\partial_\tau Z_\tau(\gamma)/|\partial_\tau Z_\tau(\gamma)|$
- **NormalizeApprox**[*z\_*, *eps\_*]: normalizes  $z \in \mathbb{C}$  to roughly unit length for large  $z$ , but behaves smoothly near zero.
- **TotalChargech2**[*Tree\_*]: gives the total charge vector  $[r, d, ch_2]$  of a given tree (nested list).
- **TotalChargechi**[*Tree\_*]: gives the total charge vector  $[r, d, \chi]$  of a given tree (nested list).
- **ConifoldRay**[*init\_*, *psi\_*, *homshift\_*]: gives a function  $f : a \in [0, 1] \rightarrow \mathbb{H}$  parametrizing the ray starting at the rational number  $init = \frac{p}{q}$  (with  $q \not\equiv 0 \pmod{3}$ )
- **RayCh**[*psi\_*]: gives an initial ray starting at 0, namely a function  $f : [0, 1] \rightarrow \mathbb{H}$  starting (close to) 0 and following a ray where  $Z_\tau([1, 0, 0]) = -T_D$  has phase  $\psi + \frac{\pi}{2} \pmod{\pi}$ . Shifting  $\psi$  by  $2\pi$  gives a different ray, corresponding to a homological shift by 2. Values are cached.
- **RayGeneralch2**[*psi\_*, *tauexpr\_*, *start\_*]: gives a function  $a \in [0, 1] \rightarrow \mathbb{H}$  parametrizing the ray where  $Z_\tau(\gamma)$  has phase  $\psi + \frac{\pi}{2} \pmod{\pi}$ . The starting point is obtained using **FindRoot**[ $\dots, Z_{tauexpr}(\gamma), \dots, start$ ], see documentation of **FindRoot**.
- **RayFromInfinity**[{*r\_*, *d\_*, *chi\_*} *psi\_*]: gives a function  $f : a \in [0, 1] \rightarrow \mathbb{H}$  parametrizing the ray of phase  $\psi$  starting from the large volume limit
- **StabilityWall**[*Tree\_*, *tauinit\_*, *tangent\_*, {*ainit\_*, *amin\_*, *amax\_*}]: gives a function  $f : a \in [0, 1] \rightarrow \mathbb{H}$  parametrizing the stability wall for the last fusion of the tree. The tree can also be a pair of charges. The *tauinit* is used as a starting point of **FindRoot** along a vertical line. The last argument can be omitted and defaults to  $\{-2, 2\}$ ; it is an interval around the starting point 0, and can be used to restrict the stability wall to only one side of *tauinit*.
- **TreeToRays**[*Tree\_*, *psi\_*]: gives the (flat) list of rays (functions  $f : [0, 1] \rightarrow \mathbb{H}$  where  $Z_\tau(\gamma_e)$  has phase  $\psi + \frac{\pi}{2} \pmod{\pi}$  along each edge with charge  $\gamma_e$ . The tree is given as a nested list of initial objects of the form  $kCh[p/q][n]$  with  $k, p, q, n$  integers.
- **TreeToRaysPlot**[*Tree\_*, *psi\_*, *plotoptions\_*]: Plots the rays produced by **TreeToRays**[*Tree*, *psi*] with the given plot options.

## 8. Higgs branch formula

In addition, the package includes some routines from the Mathematica package `CoulombHiggs.m` [2], mainly for the purpose of evaluating the indices  $K_m(p, q)$  of the Kronecker quiver. The names of the routines are prefaced by P2 to avoid clash.

- `P2HiggsBranchFormula[Mat_, Cvec_, Nvec_ ]`: computes the refined index of a quiver with DSZ products  $\alpha_{ij} = \text{Mat}[[i, j]]$ , dimension vector  $N_i = \text{Nvec}[[i]]$ , FI parameters  $\zeta_i = \text{Cvec}[[i]]$ , using Reineke's formula. It is assumed, but not checked, that the quiver has no oriented loop;
- `P2RationalInvariant[Mat_, Cvec_, Nvec_, y_ ]`: computes the refined rational index of a quiver with DSZ matrix  $\alpha_{ij} = \text{Mat}[[i, j]]$ , FI parameters  $\zeta_i = \text{Cvec}[[i]]$ , dimension vector  $N_i = \text{Nvec}[[i]]$ , using Reineke's formula
- `P2StackInvariant[Mat_, Cvec_, Nvec_, y_ ]`: computes the stacky invariant of a quiver with DSZ matrix  $\alpha_{ij} = \text{Mat}[[i, j]]$ , FI parameters  $\zeta_i = \text{Cvec}[[i]]$ , dimension vector  $N_i = \text{Nvec}[[i]]$ , using Reineke's formula
- `P2BinarySplits[Nvec_ ]`: gives the list of dimension vectors  $\gamma_L$  less than  $\gamma$ , quotiented by the equivalence relation  $\gamma_L \rightarrow \gamma - \gamma_L$ .
- `P2ListAllPartitions[gam_ ]`: returns the list of unordered partitions of the positive integer vector  $\gamma$  as a sum of positive, non-zero integer vectors  $\alpha_i$ ;
- `P2QDeformedFactorial[n_, y_ ]`: computes the  $q$ -deformed factorial  $[n, y]!$

## References

- [1] P. Bousseau, "Scattering diagrams, stability conditions, and coherent sheaves on  $\mathbb{P}^2$ ," 1909.02985.
- [2] "CoulombHiggs.m, a Mathematica package for computing quiver invariants." available from <https://github.com/bpioline/CoulombHiggs>.