

# PARMEC USER MANUAL

March 14, 2017

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# Chapter 1

## Running

PARMEC is a command line program. Typical usage:

1. Place PARMEC in a globally accessible path (e.g. suitably extend the PATH variable on a unix system).
2. Create a directory where your input file and output files will be stored (e.g. *mkdir test*).
3. Edit your Python input file in this directory (e.g. *test.py*); Chapter 3 documents all input commands.
4. Run PARMEC (e.g. *PARMEC path/to/test/test.py*).
5. Time histories can be generated during analysis using the HISTORY command; see Section 3.16.
6. Upon termination a \*.dump file and/or a \*.vtk.\* file(s) is/are created in the same directory (e.g. *path/to/test/test.dump*) their format is documented in Chapter 4.
7. The output files can be viewed with OVITO, ParaView or VisIt, as documented in Chapter 5.

## Chapter 2

# Implementation status

Individual features of parmec which are not implemented yet are marked as (under development). Relatively complex features that have seen little testing are marked as (experimental). Table 2.1 summarizes current limitations of automatic contact detection.

	SPHERE	MESH	OBSTACLE
SPHERE	OK	N/A	OK
MESH		N/A	N/A
OBSTACLE			N/A

Table 2.1: Current limitations of automatic contact detection.

## Chapter 3

# Input commands

PARMEC input language extends Python. Subroutines related to input processing are listed below.

### 3.1 RESET

Erase all data.

**RESET** ()

### 3.2 TSERIES

Create time series: a linear spline based on series of 2-points.

**tmsnum** = **TSERIES** (points)

- **tmsnum** - time series number
- **points** - a constant  $v0$ , or a list  $[t0, v0, t1, v1, \dots]$  or  $[[t0, v0], [t1, v1], \dots]$  of points (where  $t_i < t_j$ , when  $i < j$ ), or a path to a file storing pairs of times and values in format:

```
# comment 1 ...
# comment 2 ...
t0 v0
t1 v1
# comment 3 ...
t2 v2
...
```

### 3.3 MATERIAL

Create material.

**matnum** = **MATERIAL** (density, young, poisson)

- **matnum** - material number
- **density** - mass density

- **young** - Young modulus
- **poisson** - Poisson ratio

### 3.4 SPHERE

Create a spherical particle.

**parnum = SPHERE (center, radius, material, color)**

- **parnum** - particle number
- **center** - tuple  $(x, y, z)$  defining the center
- **radius** - radius
- **material** - material number
- **color** - positive integer surface color

### 3.5 MESH

Create a meshed particle.

**parnum = MESH (nodes, elements, material, colors)**

- **parnum** - particle number
- **nodes** - list of nodes:  $[x0, y0, z0, x1, y1, z1, \dots]$
- **elements** - list of elements:  $[e1, n1, n2, \dots, ne1, me1, e2, n1, n2, \dots, ne2, me2, \dots]$ , where  $e1$  is the number of nodes of the first element,  $n1, n2, \dots, ne1$  enumerate the element nodes, and  $me1$  is the material number. Similarly for the second and all remaining elements. Supported numbers of nodes per element are 4, 5, 6, and 8 for respectively *tetrahedron*, *pyramid*, *wedge*, and *hexahedron*, cf. Figure 3.1.
- **material** - material number
- **colors** - list of positive integer face colors:  $[gcolor, f1, n1, n2, \dots, nf1, c1, f2, n1, n2, \dots, nf2, c2, \dots]$ , where  $gcolor$  is the global color for all not specified faces,  $f1$  is the number of nodes in the first specified face,  $n1, n2, \dots, nf1$  enumerate the face nodes, and  $c1$  is the surface color of that face. Similarly for the second and all remaining faces. If only the global color is required, it can be passed as  $[gcolor]$  or as  $gcolor$  alone.

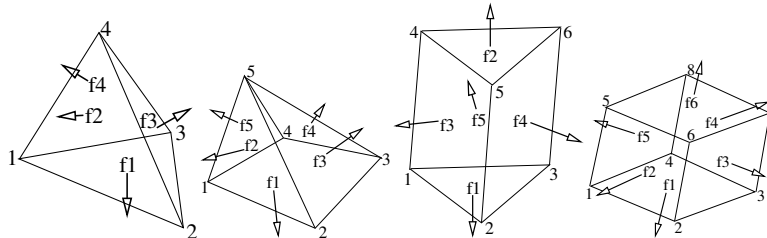


Figure 3.1: Element types.

### 3.6 ANALYTICAL

Create an analytical particle. Analytical particles have no shapes and are not involved in contact.

**parnum** = ANALYTICAL ( | **inertia**, **mass**, **rotation**, **position**, **material**, **particle**)

Note, that all parameters are optional.

- **parnum** - particle number
- **inertia** - inertia tensor passed as a list [ $I_{xx}$ ,  $I_{yy}$ ,  $I_{zz}$ ,  $I_{xy}$ ,  $I_{xz}$ ,  $I_{yz}$ ]; optional, if **particle** parameter is used; default [1, 1, 1, 0, 0, 0]
- **mass** - scalar mass; optional, if **particle** parameter is used; default 1
- **rotation** - optional orientation matrix passed as a list [ $e1x$ ,  $e1y$ ,  $e1z$ ,  $e2x$ ,  $e2y$ ,  $e2z$ ,  $e3x$ ,  $e3y$ ,  $e3z$ ], where vectors  $e1$ ,  $e2$ ,  $e3$  are orthonormal; default [1, 0, 0, 0, 1, 0, 0, 0, 1]
- **position** - optional position vector passed as a tuple ( $x$ ,  $y$ ,  $z$ ); default (0, 0, 0)
- **material** - material number; default 0
- **particle** - optional; if specified, an existing particle is converted into an analytical particle; its properties are inherited or overwritten, depending on whether any of the **inertia**, **mass**, **rotation**, **position** parameters are used; if initially specified, particle shape is inherited and its animated motion is included into the results

### 3.7 OBSTACLE

Create an obstacle.

**OBSTACLE** (**triangles**, **color** | **point**, **linear**, **angular**)

- **triangles** - list of triangle tuples [( $t1x1$ ,  $t1y1$ ,  $t1z1$ ,  $t1x2$ ,  $t1y2$ ,  $t1z2$ ,  $t1x3$ ,  $t1y3$ ,  $t1z3$ ), ( $t2x1$ ,  $t2y1$ ,  $t2z1$ ,  $t2x2$ ,  $t2y2$ ,  $t2z2$ ,  $t2x3$ ,  $t2y3$ ,  $t2z3$ ), ...] defining the obstacle
- **color** - positive integer surface color or a list [ $color1$ ,  $color2$ , ...] of colors for each individual triangle
- **point** - spatial reference point
- **linear** - linear velocity history callback: ( $v_x, v_y, v_z$ ) = **linear** ( $t$ )
- **angular** - spatial angular velocity history callback: ( $\omega_x, \omega_y, \omega_z$ ) = **angular** ( $t$ )

### 3.8 SPRING

Create a translational spring constraint. The applied force formula reads

$$\text{force}(t) = \text{direction}(t) \cdot [\text{spring}(\text{stroke}(t)) + \text{dashpot}(\text{velocity}(t)) \cdot |\text{sign}(\text{spring}(\text{stroke}(t)))|]$$

where

$$\text{direction}(t) = (\text{point2}(t) - \text{point1}(t)) / |\text{point2}(t) - \text{point1}(t)| \text{ or constant } (d_x, d_y, d_z) \text{ or tangent}$$

$$\text{stroke}(t) = \text{direction}(t) \cdot [\text{point2}(t) - \text{point1}(t)] - \text{direction}(0) \cdot [\text{point2}(0) - \text{point1}(0)]$$

$$\text{velocity}(t) = \text{direction}(t) \cdot \frac{d}{dt} [\text{point2}(t) - \text{point1}(t)]$$

$$\text{sign}(x) = \begin{cases} -1 & \text{if } x < 0 \\ 0 & \text{if } x = 0 \\ 1 & \text{if } x > 0 \end{cases}$$

The spring (stroke) and dashpot (velocity) relationships are defined by means of lookup tables; force ( $t$ ) is applied at point2 ( $t$ ), and  $-\text{force}(t)$  is applied at point1 ( $t$ ); dashpot force is not applied when spring force is zero.

**sprnum = SPRING (part1, point1, part2, point2, spring | dashpot, direction, planar, unload, ylim)**

- **sprnum** - spring number
- **part1** - first particle number
- **point1** - tuple  $(x, y, z)$  defining a point moving with the first particle
- **part2** - second particle number;  $-1$  can be used to indicate a single-particle constraint
- **point2** - tuple  $(x, y, z)$  defining a second point, either moving with the second particle, or a spatial point
- **spring** - spring force lookup table  $[\text{stroke}_1, \text{force}_1, \text{stroke}_2, \text{force}_2, \dots, \text{stroke}_n, \text{force}_n]$ ; used for both loading and unloading when the **unload** table and the **yield** limits are not given
- **dashpot** - optional dashpot force lookup table  $[\text{velocity}_1, \text{force}_1, \text{velocity}_2, \text{force}_2, \dots, \text{velocity}_m, \text{force}_m]$ ; default:  $[-\infty, 0, +\infty, 0]$
- **direction** - optional constant direction  $(d_x, d_y, d_z)$
- **planar** - optional planar spring flag; when 'ON' spring direction

$$(\text{point2}(t) - \text{point1}(t)) / |\text{point2}(t) - \text{point1}(t)|$$

is projected onto a plane orthogonal to  $(d_x, d_y, d_z)$ ; default: 'OFF'

- **unload** - spring unloading lookup table  $[\text{stroke}_1, \text{force}_1, \text{stroke}_2, \text{force}_2, \dots, \text{stroke}_n, \text{force}_n]$ ; must be monotonically increasing
- **ylim** - tuple  $(f_{yc}, f_{yt})$  defining the compression,  $f_{yc} < 0$ , and tension,  $f_{yt} > 0$ , yield limits; the unloading curve begins to be used once either of these limits is crossed; default:  $(0, 0)$

### 3.9 GRANULAR

Define surface pairing for the granular contact interaction model.

**GRANULAR (color1, color2, spring | damper, friction, rolling, drilling, kskn)**

- **color1** - first color (positive, or color1 = 0 and color2 = 0 to redefine default parameters)
- **color2** - second color (positive, or color1 = 0 and color2 = 0 to redefine default parameters)
- **spring** - normal spring constant
- **damper** - optional normal damping ratio; default: 1.0



- **friction** - optional Coulomb's friction coefficient; default: 0.0; tuple  $(\mu_s, \mu_d)$  can be used to specify respectively static and dynamic friction coefficients; (experimental)
- **rolling** - optional rolling friction coefficient; default: 0.0; (under development)
- **drilling** - optional drilling friction coefficient; default: 0.0; (under development)
- **kskn** - optional ratio of normal to tangential spring and dashpot parameters; default: 0.5

### 3.10 CONSTRAIN

Constrain particle motion.

#### CONSTRAIN (parnum | linear, angular)

- **parnum** - particle number
- **linear** - list  $[x_1, y_1, z_1]$ ,  $[x_1, y_1, z_1, x_2, y_2, z_2]$ , or  $[x_1, y_1, z_1, x_2, y_2, z_2, x_3, y_3, z_3]$  defining directions of constrained linear motion; default:  $[0, 0, 0]$
- **angular** - list  $[x_1, y_1, z_1]$ ,  $[x_1, y_1, z_1, x_2, y_2, z_2]$ , or  $[x_1, y_1, z_1, x_2, y_2, z_2, x_3, y_3, z_3]$  defining directions of constrained spatial rotation; default:  $[0, 0, 0]$

### 3.11 PRESCRIBE

Prescribe particle motion. Prescribed motion overwrites this resulting from dynamics and constraints.

#### PRESCRIBE (parnum | linear, angular, kind)

- **parnum** - particle number
- **linear** - a tuple  $(i, j, k)$  of TSERIES numbers, or a callback:  $(v_x, v_y, v_z) = \mathbf{linear}(t)$ , defining linear velocity or acceleration history; default: *not prescribed*
- **angular** - a tuple  $(i, j, k)$  of TSERIES numbers, or a callback:  $(\omega_x, \omega_y, \omega_z) = \mathbf{angular}(t)$ , defining spatial angular velocity or acceleration history; default: *not prescribed*
- **kind** - string 'vv', 'va', 'av', or 'aa' indicating interpretation of respectively **linear** and **angular** time histories as either velocity or acceleration; default: 'vv'

### 3.12 VELOCITY

Set particle velocity.

#### VELOCITY (parnum | linear, angular)

- **parnum** - particle number
- **linear** - linear velocity tuple  $(v_x, v_y, v_z)$ ; default:  $(0, 0, 0)$  at  $t = 0$
- **angular** - angular velocity tuple  $(\omega_x, \omega_y, \omega_z)$ ; default:  $(0, 0, 0)$  at  $t = 0$

### 3.13 GRAVITY

Set gravity.

#### GRAVITY (**gx**, **gy**, **gz**)

- **gx** - constant  $x$  float number, or callback **gx**( $t$ ), or TSERIES number
- **gy** - constant  $y$  float number, or callback **gy**( $t$ ), or TSERIES number
- **gz** - constant  $z$  float number, or callback **gz**( $t$ ), or TSERIES number

### 3.14 DAMPING

Set global damping, applied as

$$\text{force} = -m \begin{bmatrix} -d_{vx}v_x \\ -d_{vy}v_y \\ -d_{vz}v_z \end{bmatrix}, \text{ torque} = -\mathbf{\Lambda}\mathbf{J}\mathbf{\Lambda}^T \begin{bmatrix} -d_{\omega x}\omega_x \\ -d_{\omega y}\omega_y \\ -d_{\omega z}\omega_z \end{bmatrix}$$

where  $m$  is scalar mass,  $v$  is linear velocity,  $\mathbf{\Lambda}$  is the rotation matrix,  $\mathbf{J}$  is the referential inertia matrix, and  $\omega$  is spatial angular velocity.

#### DAMPING (**linear**, **angular**)

- **linear** - linear damping curve callback:  $(d_{vx}, d_{vy}, d_{vz}) = \mathbf{linear}(t)$
- **angular** - angular damping curve callback:  $(d_{\omega x}, d_{\omega y}, d_{\omega z}) = \mathbf{angular}(t)$

### 3.15 CRITICAL

Estimate critical time step.

#### **h** = CRITICAL ()

- **h** - critical time step

### 3.16 HISTORY

Before running a simulation, request time history output.

#### **list** = HISTORY (**entity** | **source**, **point**)

- **list** - output time history list (empty upon initial request, populated during simulation)
- **entity** - entity name; global entities: (output time) 'TIME'; particle entities: (position) 'PX', 'PY', 'PZ', '|P|', (displacement) 'DX', 'DY', 'DZ', '|D|', (linear velocity) 'VX', 'VY', 'VZ', '|V|', (angular velocity) 'OX', 'OY', 'OZ', '|O|', (body force) 'FX', 'FY', 'FZ', '|F|', (body torque) 'TX', 'TY', 'TZ', '|T|'; spring entities: (spring stroke) 'STROKE', (spring total force) 'STF', (spring force without damping) 'SF';

- **source** - particle number  $i$ , or a list of particle numbers  $[i, j, \dots]$ , or a spatial sphere defined as tuple  $(x, y, z, r)$ , or a spatial box defined as tuple  $(x_{\min}, y_{\min}, z_{\min}, x_{\max}, y_{\max}, z_{\max})$ ; in case of a list of particle numbers the output entity is averaged over the set of particles; in case of a spatial sphere or box the output entity is averaged over the set of particles passing through it; default: 0 (useful when entity is 'TIME'); spring number or a list of numbers can be used as a source in case of spring entities
- **point** - optional referential point used in case of a single particle source; default: particle mass centre

### 3.17 OUTPUT

Before running a simulation, define scalar and/or vector entities included into the output file(s). PARMEC outputs:

- \*.dump files for spherical particles
- \*0.vtk.\* **and/or** (\*0.h5, \*0.xmf) files for obstacles and mesh based particles **not** specified as **a subset** in the OUTPUT command
- \*1.vtk.\*, \*2.vtk.\*, ... **and/or** (\*1.h5, \*1.xmf, \*2.h5, \*2.xmf, ...) files for mesh based particles specified as subsets, where numbers 1, 2, ... match consecutive OUTPUT calls
- \*0rb.vtk.\* **and/or** (\*0rb.h5, \*0rb.xmf) for rigid body data of particles **not** specified as **a subset** in the OUTPUT command
- \*1rb.vtk.\*, \*2rb.vtk.\*, ... **and/or** (\*1rb.h5, \*1rb.xmf, \*2rb.h5, \*2rb.xmf, ...) files for rigid body data of particles specified as subsets, where numbers 1, 2, ... match consecutive OUTPUT calls
- \*0cd.vtk.\* **and/or** (\*0cd.h5, \*0cd.xmf) for contact data including particles **not** specified as **a subset** in the OUTPUT command
- \*1cd.vtk.\*, \*2cd.vtk.\*, ... **and/or** (\*1cd.h5, \*1cd.xmf, \*2cd.h5, \*2cd.xmf, ...) files for contact data including particles specified as subsets, where numbers 1, 2, ... match consecutive OUTPUT calls
- \*0sd.vtk.\* **and/or** (\*0sd.h5, \*0sd.xmf) for spring data including particles **not** specified as **a subset** in the OUTPUT command
- \*1sd.vtk.\*, \*2sd.vtk.\*, ... **and/or** (\*1sd.h5, \*1sd.xmf, \*2sd.h5, \*2sd.xmf, ...) files for spring data including particles specified as subsets, where numbers 1, 2, ... match consecutive OUTPUT calls

#### OUTPUT (entities | subset, mode, format)

- **entities** - list of output entities; default: ['NUMBER', 'COLOR', 'DISPL', 'ORIENT', 'LINVEL', 'ANGVEL', 'FORCE', 'TORQUE', 'F', 'FN', 'FT', 'SF', 'AREA', 'PAIR'] where:
  - 'NUMBER' - scalar field of particle numbers (modes: 'SPH', 'MESH', 'RB'), or scalar field of spring numbers (modes: 'SD')
  - 'COLOR' - scalar field of surface colors (modes: 'SPH', 'MESH'), or 2-component vector field of contact surface colors (modes: 'CD')
  - 'DISPL' - 3-component vector field of displacements (modes: 'SPH', 'MESH', 'RB'), or scalar field of contact depths (modes: 'CD'), or scalar field of spring strokes (modes: 'SD')
  - 'ORIENT' - three 3-component vector fields representing columns of rigid rotation matrix (orientation vectors) (modes: 'RB'), or 3-component vector field of spring orientations (modes: 'SD')
  - 'LINVEL' - 3-component vector field of linear velocity (modes: 'SPH', 'MESH', 'RB')

- 'ANGVEL' - 3-component vector field of (spatially constant) angular velocity (modes: 'SPH', 'MESH', 'RB')
- 'FORCE' - 3-component vector field of (spatially constant) total body force (modes: 'SPH', 'MESH', 'RB')
- 'TORQUE' - 3-component vector field of (spatially constant) total body torque (modes: 'SPH', 'MESH', 'RB')
- 'F' - 3-component vector field of total contact forces (modes: 'CD'), or scalar field of total spring forces (modes: 'SD')
- 'FN' - 3-component vector field of normal contact forces (modes: 'CD')
- 'FT' - 3-component vector field of tangential contact forces (modes: 'CD')
- 'SF' - scalar field of spring force magnitude, without dashpot contribution (modes: 'CD', 'SD')
- 'AREA' - scalar field of contact area (modes: 'CD')
- 'PAIR' - 2-component vector field of particle pair numbers (modes: 'CD', 'SD')
- **subset** - optional particle number  $i$ , or a list of particle numbers  $[i, j, \dots]$ , to which this specification is narrowed down
- **mode** - optional output mode or list of output modes: 'SPH' for sphere output, 'MESH' for mesh output, 'RB' for rigid body output, 'CD' for contact data output, 'SD' for spring data output; default: ['SPH', 'MESH', 'RB', 'CD', 'SD']
- **format** - optional output format, e.g. 'VTK' or 'XDMF', or list ['VTK', 'XDMF'], where 'VTK' is the text based legacy VTK format, 'XDMF' is the HDF5/XML based XDMF format; default: 'XDMF'

### 3.18 DEM

Run DEM simulation.

**t = DEM (duration, step | interval, prefix, adaptive)**

- **t** - simulation runtime in seconds
- **duration** - simulation duration
- **step** - time step; initial if **adaptive** is used or constant otherwise
- **interval** - output interval (default: time step); tuple  $(dt_{\text{files}}, dt_{\text{history}})$  can be used to indicate different output frequencies of output files and time histories, respectively; callback functions can also be used, e.g.  $dt_{\text{files}} = dt_{\text{files}}(t)$  and  $dt_{\text{history}} = dt_{\text{history}}(t)$  prescribing variable interval frequencies, depending on current time;
- **prefix** - output file name prefix (default: input file name without the ".py" extension)
- **adaptive** - adaptive time step reduction factor; zero turns off adaptive time stepping, values  $> 0.0$  and  $\leq 1.0$  turn it on; default: 0.0 (experimental)

## Chapter 4

# Output files

(Under development)

## Chapter 5

# Output viewers

(Under development)