
ParaView Users Guide Documentation

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ParaView Developers

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PARAVIEW USER'S GUIDE

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This guide can be split into two volumes. The **User's Guide** covers various aspects of data analysis and visualization with **ParaView**.

Reference Manual provides details on various components in the UI and the scripting API.

PARAVIEW USER'S GUIDE

1.1 Introduction to ParaView

1.1.1 Introduction

ParaView is an open-source, multi-platform scientific data analysis and visualization tool that enables analysis and visualization of extremely large datasets. **ParaView** is both a general purpose, end-user application with a distributed architecture that can be seamlessly leveraged by your desktop or other remote parallel computing resources and an extensible framework with a collection of tools and libraries for various applications including scripting (using Python), web visualization (through ParaViewWeb), or in-situ analysis (with Catalyst).

ParaView leverages parallel data processing and rendering to enable interactive visualization for extremely large datasets. It also includes support for large displays including tiled displays and immersive 3D displays with head tracking and wand control capabilities.

ParaView also supports scripting and batch processing using Python. Using included Python modules, you can write scripts that can perform almost all the functionality exposed by the interactive application and much more.

ParaView is open-source (BSD licensed, commercial software friendly). As with any successful open-source project, **ParaView** is supported by an active user and developer community. Contributions to both the code and this user's manual that help make the tool and the documentation better are always welcome.

Did you know?

The **ParaView** project started in 2000 as a collaborative effort between Kitware Inc. and LANL (Los Alamos National Labs). The initial funding was provided by a three year contract with the US Department of Energy ASCI Views program. The first public release, **ParaView** 0.6, was announced in October 2002.

Independent of **ParaView**, Kitware started developing a web-based visualization system in December 2001. This project was funded by Phase I and II SBIRs from the US ARL (Arm Research Laboratory) and eventually became the PVEE (ParaView Enterprise Edition). PVEE significantly contributed to the development of **ParaView**'s client/server architecture. PVEE was the precursor to ParaViewWeb, a modern web visualization solution based on **ParaView**.

Since the project began, Kitware has successfully collaborated with Sandia, LANL, the ARL, and various other academic and government institutions to continue development. Today, the project is still going strong!

In September 2005, Kitware, Sandia National Labs and CSimSoft started the development of **ParaView** 3.0. This was a major effort focused on rewriting the user interface to be more user friendly and on developing a quantitative analysis framework. **ParaView** 3.0 was released in May 2007.

In this guide

This user's manual is designed as a guide for using the **ParaView** application. It is geared toward users who have a general understanding of common data visualization techniques. For scripting, a working knowledge of the Python language is assumed. If you are new to Python, there are several tutorials and guides for getting started that are available on the Internet.

Did You Know?

In this guide, we will periodically use these **Did you know?** boxes to provide additional information related to the topic at hand.

Common Errors

Common Errors blocks are used to highlight some of the common problems or complications you may run into when dealing with the topic of discussion.

This guide can be split into two volumes. [Section 1.1](#) to [Section 1.8](#) can be considered the user's guide, where various aspects of data analysis and visualization with **ParaView** are covered. [Section 2.1](#) to [Section 2.11](#) in Reference manual part provide details on various components in the UI and the scripting API.

Getting help

This guide tries to cover most of the commonly used functionality in **ParaView**. **ParaView**'s flexible, pipeline-based architecture opens up numerous possibilities. If you find yourself looking for some feature not covered in this guide, refer to the Wiki pages (<http://paraview.org/Wiki/ParaView>) or to the forum (<https://discourse.paraview.org/categories>) specially the **|FAQ|** and the **|Tips and Tricks|** categories. Also feel free to ask about it under the relevant **|Support|** category.

Getting the software

ParaView is open source. The complete source code for all the functionality discussed in The **ParaView** Guide can be downloaded from the **ParaView** website <http://www.paraview.org>. We also provide binaries for the major platforms: Linux, Mac OS X, and Windows. You can get the source files and binaries for the official releases, as well as follow **ParaView**'s active development, by downloading the nightly builds.

Providing details of how to build **ParaView** using the source files is beyond the scope of this guide. Refer to the **ParaView** gitlab (<https://gitlab.kitware.com/paraview/paraview/-/blob/master/Documentation/dev/build.md>) for more information.

1.1.2 Basics of visualization in ParaView

Visualization is the process of converting raw data into images and renderings to gain a better cognitive understanding of the data. **ParaView** uses VTK, the Visualization Toolkit, to provide the backbone for visualization and data processing.

The VTK model is based on the data-flow paradigm. In this paradigm, data flows through the system being transformed at each step by modules known as algorithms. Algorithms could be common operations such as clipping, slicing, or generating contours from the data, or they could be computing derived quantities, etc. Algorithms have input ports through which they take in data and output ports through which they produce output. You need producers that ingest data into the system. These are simply algorithms that do not have an input port but have one or more output ports. They are called *sources*. Readers that read data from files are examples of such sources. Additionally, there are algorithms

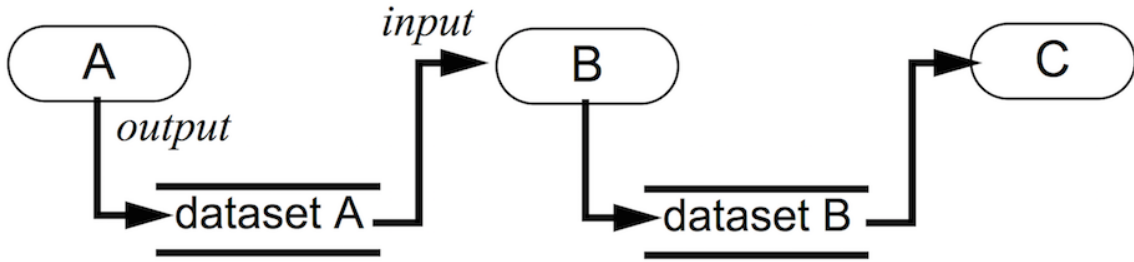


Fig. 1.1: Visualization model: Process objects A, B, and C input and/or output one or more data objects. Data objects represent and provide access to data; process objects operate on the data. Objects A, B, and C are source, filter, and mapper objects, respectively. [SML96]

that transform the data into graphics primitives so that they can be rendered on a computer screen or saved to disk in another file. These algorithms, which have one or more input ports but do not have output ports, are called *sinks*. Intermediate algorithms with input ports and output ports are called *filters*. Together, sources, filters, and sinks provide a flexible infrastructure wherein you can create complex processing pipelines by simply connecting algorithms to perform arbitrarily complex tasks.

For more information on VTK's programming model, refer to [SML96].

This way of looking at the visualization pipeline is at the core of **ParaView**'s work flow: You bring your data into the system by creating a reader – the source. You then apply filters to either extract information (e.g., iso-contours) and render the results in a view or to save the data to disk using writers – the sinks.

ParaView includes readers for a multitude of file formats typically used in the computational science world. To efficiently represent data from various fields with varying characteristics, VTK provides a rich data model that **ParaView** uses. The data model can be thought of simply as ways of representing data in memory. We will cover the different data types in more detail in Section 1.3.1. Readers produce a data type suitable for representing the information the files contain. Based on the data type, **ParaView** allows you to create and apply filters to transform the data. You can also show the data in a view to produce images or renderings. Just as there are several types of filters, each performing different operations and types of processing, there are several kinds of views for generating various types of renderings including 3D surface views, 2D bar and line views, parallel coordinate views, etc.

Did You Know?

The Visualization Toolkit (VTK) is an open-source, freely available software system for 3D computer graphics, modeling, image processing, volume rendering, scientific visualization, and information visualization. VTK also includes ancillary support for 3D interaction widgets, two and three-dimensional annotation, and parallel computing.

At its core, VTK is implemented as a C++ toolkit, requiring users to build applications by combining various objects into an application. The system also supports automated wrapping of the C++ core into Python, Java, and Tcl so that VTK applications may also be written using these interpreted programming languages. VTK is used world-wide in commercial applications, research and development, and as the basis of many advanced visualization applications such as **ParaView**, VisIt, VisTrails, Slicer, MayaVi, and OsiriX.

1.1.3 ParaView executables

ParaView comes with several executables that serve different purposes.

`paraview`

This is the main **ParaView** graphical user interface (GUI). In most cases, when we refer to **ParaView**, we are indeed talking about this application. It is a Qt-based, cross-platform UI that provides access to the **ParaView** computing capabilities. Major parts of this guide are dedicated to understanding and using this application.

`pvpython`

`pvpython` is the Python interpreter that runs **ParaView**'s Python scripts. You can think of this as the equivalent of the `paraview` for scripting.

`pvbatch`

Similar to `pvpython`, `pvbatch` is also a Python interpreter that runs Python scripts for **ParaView**. The one difference is that, while `pvpython` is meant to run interactive scripts, `pvbatch` is designed for batch processing. Additionally, when running on computing resources with MPI capabilities, `pvbatch` can be run in parallel. We will cover this in more detail in Section 2.6.10.

`pvserver`

For remote visualization, this executable represents the server that does all of the data processing and, potentially, the rendering. You can make `paraview` connect to `pvserver` running remotely on an HPC resource. This allows you to build and control visualization and analysis on the HPC resource from your desktop as if you were simply processing it locally on your desktop!

`pvdataserver` and `pvrenderserver`

These can be thought of as the `pvserver` split into two separate executables: one for the data processing part, `pvdataserver`, and one for the rendering part, `pvrenderserver`. Splitting these into separate processes makes it possible to perform data processing and rendering on separate sets of nodes with appropriate computing capabilities suitable for the two tasks. Just as with `pvserver`, `paraview` can connect to a `pvdataserver`-`pvrenderserver` pair for remote visualization. Unless otherwise noted, all discussion of remote visualization or client-server visualization in this guide is applicable to both `pvserver` and `pvdataserver` - `pvrenderserver` configurations.

1.1.4 Getting started with `paraview`

`paraview` graphical user interface

`paraview` is the graphical front-end to the **ParaView** application. The UI is designed to allow you to easily create pipelines for data processing with arbitrary complexity. The UI provides panels for you to inspect and modify the pipelines, to change parameters that in turn affect the processing pipelines, to perform various data selection and inspection actions to introspect the data, and to generate renderings. We will cover various aspects of the UI for the better part of this guide.

Let's start by looking at the various components of the UI. If you run `paraview` for the first time, you will see something similar to the Fig. 1.2. The UI is comprised of menus, dockable panels, toolbars, and the viewport – the central portion of the application window.

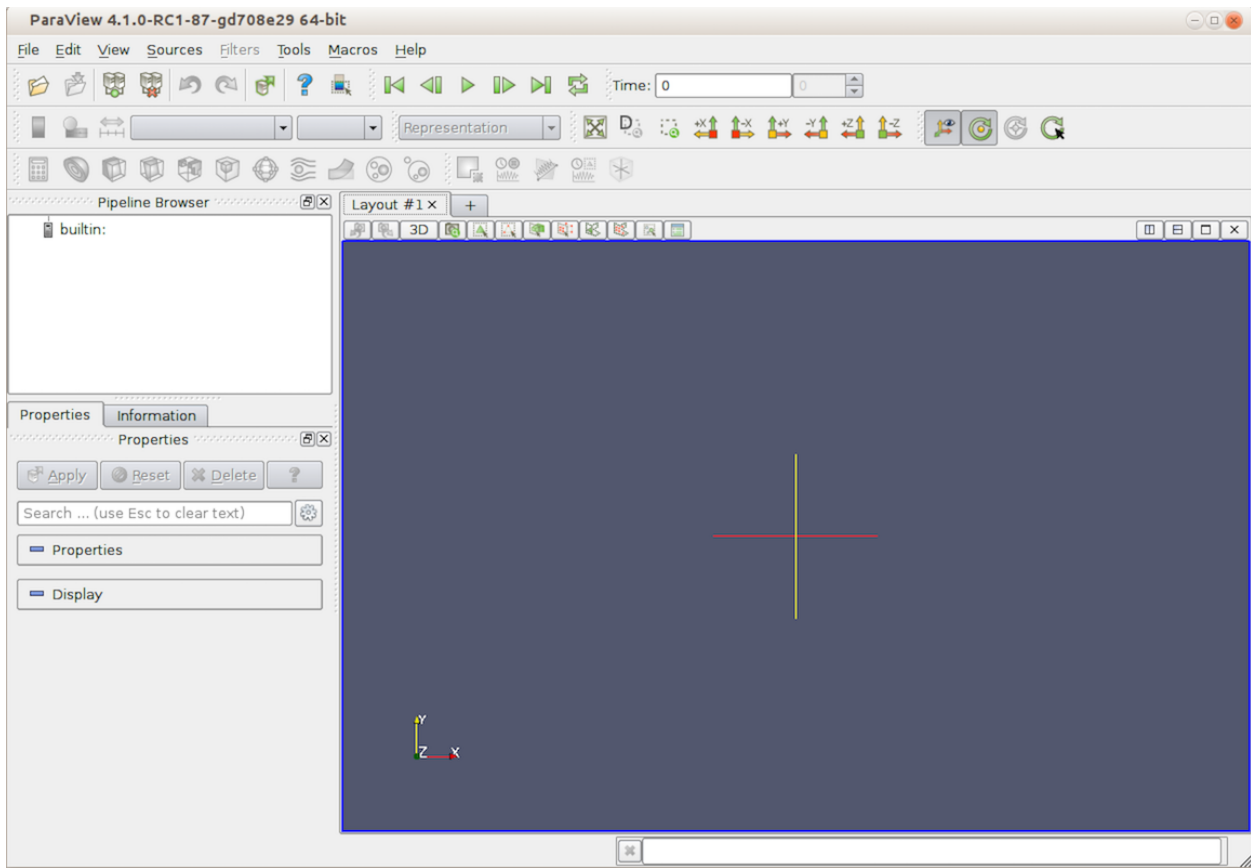


Fig. 1.2: paraview application window.

Menus provide the standard set of options typical with a desktop application including options for opening/saving files (*File* menu), for undo/redo (*Edit* menu), for the toggle panel, and for toolbar visibilities (*View* menu). Additionally, the menus provide ways to create sources that generate test datasets of various types (*Sources* menu), as well new filters for processing data (*Filters* menu). The *Tools* menu provides access to some of the advanced features in `paraview` such as managing plugins and favorites.

Panels provide you with the ability to peek into the application's state. For example, you can inspect the visualization pipeline that has been set up (*Pipeline Browser*), as well as the memory that is being used (*Memory Inspector*) and the parameters or properties for a processing module (*Properties* panel). Several of the panels also allow you to change the values that are displayed, e.g., the *Properties* panel not only shows the processing module parameters, but it also allows you to change them. Several of the panels are context sensitive. For example, the *Properties* panel changes to show the parameters from the selected module as you change the active module in the *Pipeline Browser*.

Toolbars are designed to provide quick access to common functionality. Several of the actions in the toolbar are accessible from other locations, including menus or panels. Similar to panels, some of the toolbar buttons are context sensitive and will become enabled or disabled based on the selected module or view.

The viewport or the central portion of the `paraview` window is the area where **ParaView** renders results generated from the data. The containers in which data can be rendered or shown are called *views*. You can create several different types of views, all of which are laid out in this viewport area. By default, a 3D view is created, which is one of the most commonly used views in **ParaView**.

Understanding the visualization process

To gain a better understanding of how to use the application interface, let's consider a simple example: creating a data source and applying a filter to it.

Creating a source

The visualization process in **ParaView** begins by bringing your data into the application. Section 1.2 explains how to read data from various file formats. Besides reading files to bring in data into the application, **ParaView** also provides a collection of data sources that can produce sample datasets. These are available under the *Sources* menu. To create a source, simply click on any item in the *Source* menu.

Did You Know?

As you move your cursor over the items in any menu, on most platforms (except Mac OS X), you'll see a brief description of the item in the status bar on the lower-left corner in the application window.

If you click on *Sources > Sphere*, for example, you'll create a producer algorithm that generates a spherical surface, as shown in Fig. 1.3.

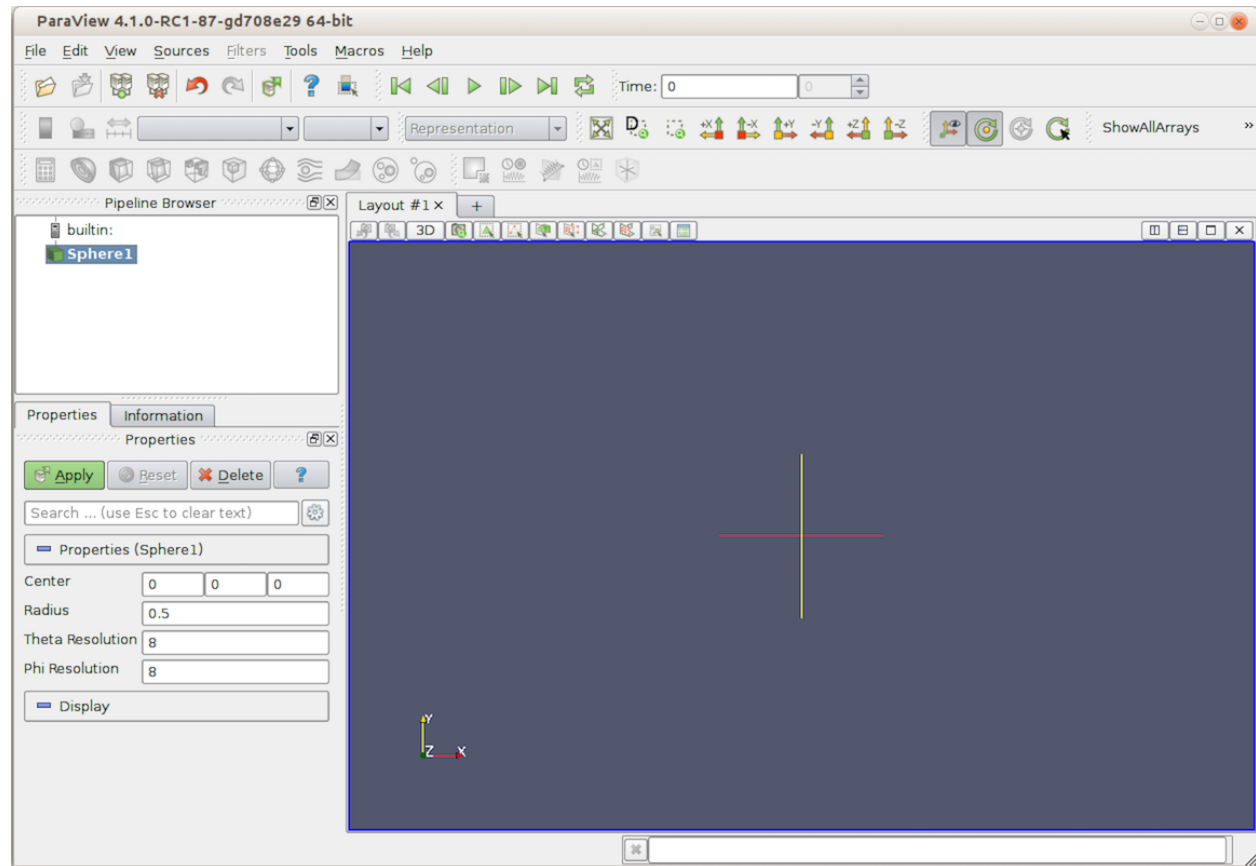


Fig. 1.3: Visualization in paraview: Step 1.

A few things to note:

1. A pipeline module is added in the Pipeline Browser panel with a name derived from the menu item, as is highlighted.
2. The Properties panel fills up with text to indicate that it's showing properties for the highlighted item (which, in this case, is Sphere1), as well as to display some widgets for parameters such as Center, Radius, etc.
3. On the Properties panel, the Apply button becomes enabled and highlighted.
4. The 3D view remains unaffected, as nothing new is shown or rendered in this view as of yet.

Let's take a closer look at what has happened. When we clicked on *Sources > Sphere*, referring to [Section 1.1.2](#), we created an instance of a source that can produce a spherical surface mesh – that's what is reflected in the Pipeline Browser. This instance receives a name, which is used by the Sphere1 and the Pipeline Browser, as well as other components of the UI, to refer to this instance of the source. Pipeline modules such as sources and filters have parameters on them that you can change that affect that module's behavior. We call them *properties*. The Properties panel shows these properties and allows you to change them. Since the ingestion of data into the system can be a time-consuming process, paraview allows you to change the properties before the module executes or performs the actual processing to ingest the data. Hence, the Apply button is highlighted to indicate that you need to accept the properties before the application will proceed. Since no data has entered the system yet, there's nothing to show. Therefore, the 3D view remains unaffected.

Let's assume we are okay with the default values for all of the properties on the Sphere1. Next, click on the Apply button.

Let's assume we are okay with the default values for all of the properties on the Sphere1. Next, click on the Apply button.

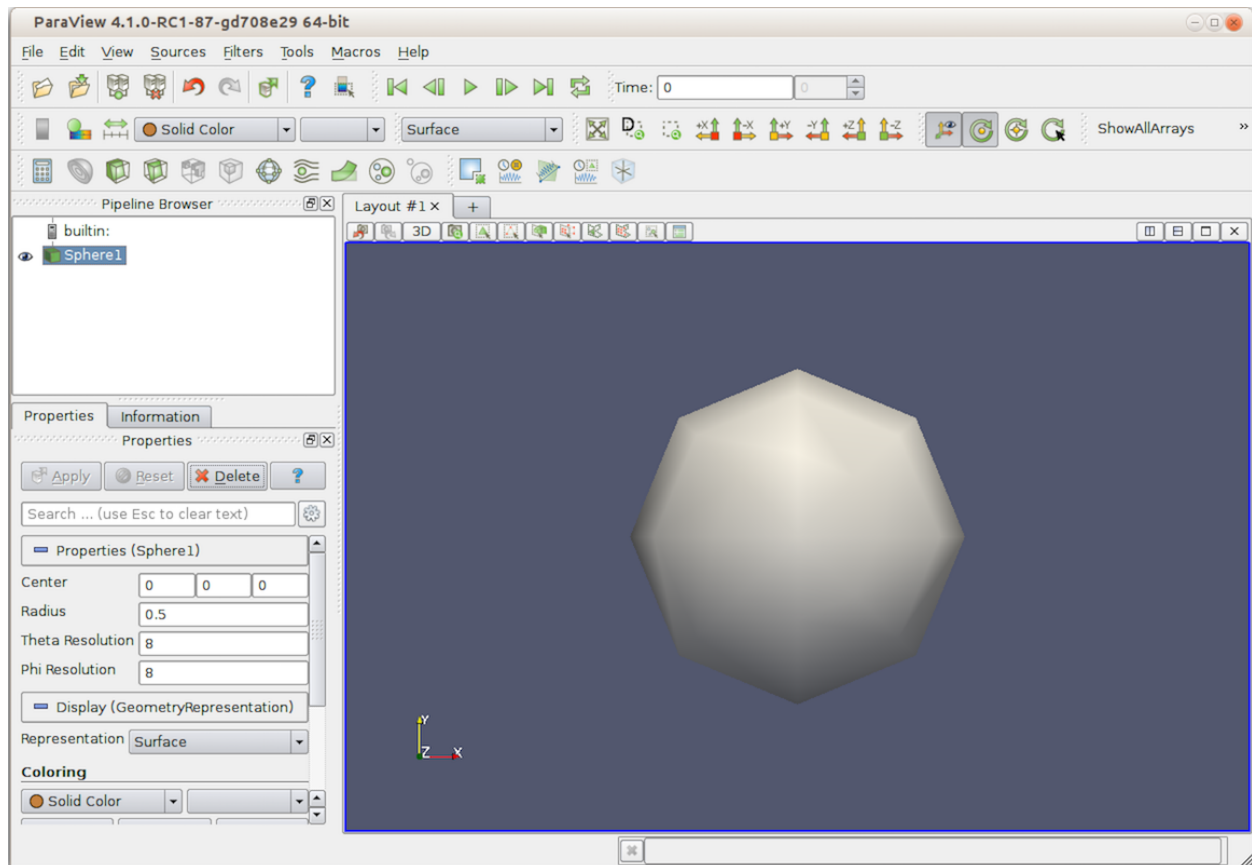


Fig. 1.4: Visualization in paraview: Step 2.

The following will ensue (Fig. 1.4):

1. The **Apply** button goes back to its old disabled/un-highlighted state.
2. A spherical surface is rendered in the 3D view.
3. The **Display** section on the **Properties** panel now shows new parameters or properties.
4. Certain toolbars update, and you can see that toolbars with text, such as **Solid Color** and **Surface** , now become enabled.

By clicking **Apply** , we told paraview to apply the properties shown on the **Properties** panel. When a new source (or filter) is applied for the first time, paraview will automatically show the data that the pipeline module produces in the current view, if possible. In this case, the sphere source produces a surface mesh, which is then shown or displayed in the 3D view.

The properties that allow you to control how the data is displayed in the view are now shown on the **Properties** panel in the **Display** section. Things such as the surface color, rendering type or representation, shading parameters, etc., are shown under this newly updated section. We will look at display properties in more detail in [Section 1.4](#).

Some of the properties that are commonly used are also duplicated in the toolbar. These properties include the data array with which the surface is colored and the representation type. These are the changes in the toolbar that allow you to quickly change some display properties.

Changing properties

If you change any of the properties on the sphere source, such as the properties under the **Properties** section on the **Properties** panel, including the **Radius** for the spherical mesh or its **Center** , the **Apply** button will be highlighted again. Once you are finished with all of the property changes, you can hit **Apply** to apply the changes. Once the changes are applied, paraview will re-execute the sphere source to produce a new mesh, as requested. It will then automatically update the view, and you will see the new result rendered.

If you change any of the display properties for the sphere source, such as the properties under the **Display** section of the **Properties** panel (including **Representation** or **Opacity**), the **Apply** button is not affected, the changes are immediately applied, and the view is updated.

The rationale behind this is that, typically, the execution of the source (or filter) is more computationally intensive than the rendering. Changing source (or filter) properties causes that algorithm to re-execute, while changing display properties, in most cases, only triggers a fresh render with an updated graphics state.

Did You Know?

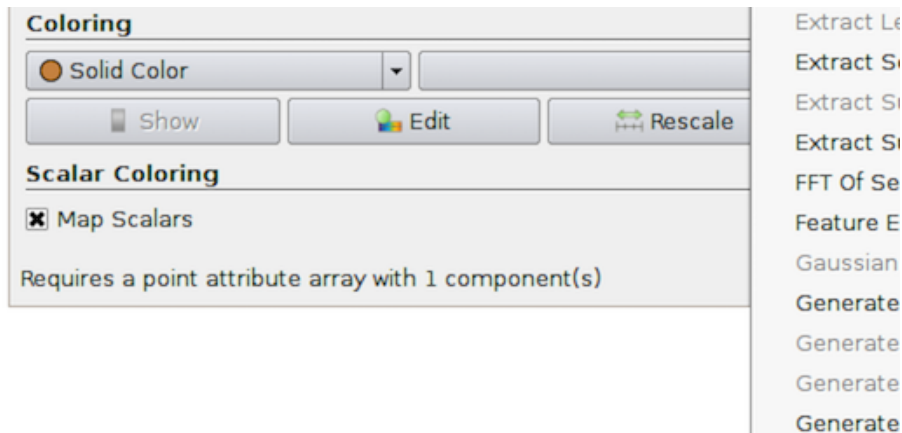
For some workflows with smaller data sizes, it may be more convenient if the **Apply** button was automatically applied even after changes are made to the pipeline module properties. You can change this from the application settings dialog, which is accessible from the *Edit > Settings* menu. The setting is called **Auto Apply** . You can also change the **Auto Apply** state using the  button from the toolbar.

Applying filters

As per the data-flow paradigm, one creates pipelines with filters to transform data. Similar to the *Sources* menu, which allows us to create new data sources, there's a *Filters* menu that provides access to the large set of filters that are available in **ParaView**. If you peruse the items in this menu, some of them will be enabled, and some of them will be disabled. Filters that can work with the data type being produced by the sphere source are enabled, while others are disabled. You can click on any of the enabled filters to create a new instance of that filter type.

Did You Know?

To figure out why a particular filter doesn't work with the current source, simply move your mouse over the disabled item in the *Filters* menu. On Linux and Windows (not OS X, however), the status bar will provide a brief explanation of why that filter is not available.



For example, if you click on *Filters > Shrink*, it will create a filter that shrinks each of the mesh cells by a fixed factor. Exactly as before, when we created the sphere source, we see that the newly-created filter is given a new name, *Shrink1*, and is highlighted in the **Pipeline Browser**. The **Properties** panel is also updated to show the properties for this new filter, and the **Apply** button is highlighted to request that we accept the properties for the filter so that it can be executed and the result can be rendered. If you click back and forth between the *Sphere1* and *Shrink1* in the **Pipeline Browser**, you'll see the **Properties** panel and toolbars update, reflecting the state of the selected pipeline module. This is an important concept in **ParaView**. There's a notion of active pipeline module, called the *active source*. Several panels, toolbars, and menus will update based on the active source.

If you click **Apply**, as was the case before, the shrink filter will be executed and the resulting dataset will be generated and shown in the 3D view. **paraview** will also automatically hide the result from the *Sphere1* so that it is not shown in the view. Otherwise, the two datasets will overlap. This is reflected by the change of state for the *eyeball* icons in the **Pipeline Browser** next to each of the pipeline modules. You can show or hide results from any pipeline module by clicking on the eyeballs.

This simple workflow forms the basis of all the data analysis and visualization in **ParaView**. The process involves creating sources and filters, changing their parameters, and showing the generated result in one or more views. In the rest of this guide, we will cover various types of filters and data processing that you can do. We will also cover different types of views that can help you produce a wide array of 2D and 3D visualizations, as well as inspect your data and drill down into it.

Common Errors

Beginners often forget to hit the **Apply** button after creating sources or filters or after changing properties. This is one

of the most common pitfalls for users new to the **ParaView** workflow.

1.1.5 Getting started with pypython

While this section refers to pypython, everything that we discuss here is applicable to pvbatch as well. Until we start looking into parallel processing, the only difference between the two executables is that pypython provides an interactive shell wherein you can type your commands, while pvbatch expects the Python script to be specified on the command line argument.

pypython scripting interface

ParaView provides a scripting interface to write scripts for performing the tasks that you could do using the GUI. The scripting interface can be accessed through Python, which is an interpreted programming language popular among the scientific community for its simplicity and its capabilities. While a working knowledge of Python will be useful for writing scripts with advanced capabilities, you should be able to follow most of the discussion in this book about **ParaView** scripting even without much Python exposure.

ParaView provides a paraview package with several Python modules that expose various functionalities. The primary scripting interface is provided by the simple module.

When you start pypython, you should see a prompt in a terminal window as follows (with some platform specific differences).

```
Python 2.7.5 (default, Sep 2 2013, 05:24:04)
[GCC 4.2.1 Compatible Apple LLVM 5.0 (clang-500.0.68)] on darwin
Type "help", "copyright", "credits" or "license" for more information
>>>
```

You can now type commands at this prompt, and **ParaView** will execute them. To bring in the **ParaView** scripting API, you first need to import the simple module from the paraview package as follows:

```
>>> from paraview.simple import *
```

Common Errors

Remember to hit the Enter or Return key after every command to execute it. Any Python interpreter will not execute the command until Enter is hit.

If the module is loaded correctly, pypython will present a prompt for the next command.

```
>>> from paraview.simple import *
>>>
```

You can consider this as in the same state as when paraview was started (with some differences that we can ignore for now). The application is ready to ingest data and start processing.

Understanding the visualization process

Let's try to understand the workflow by looking at the same use-case as we did in [Section 1.1.4](#).

Creating a source

In paraview, we created the data source by using the *Sources* menu. In the scripting environment, this maps to simply typing the name of the source to create.

```
>>> Sphere()
```

This will create the sphere source with a default set of properties. Just like with paraview, as soon as a new pipeline module is created, it becomes the *active source*.

Now, to show the active source in a view, try:

```
>>> Show()  
>>> Render()
```

The Show call will prepare the display, while the Render call will cause the rendering to occur. In addition, a new window will popup, showing the result ([Fig. 1.5](#)). This is similar to the state after hitting **Apply** in the UI.

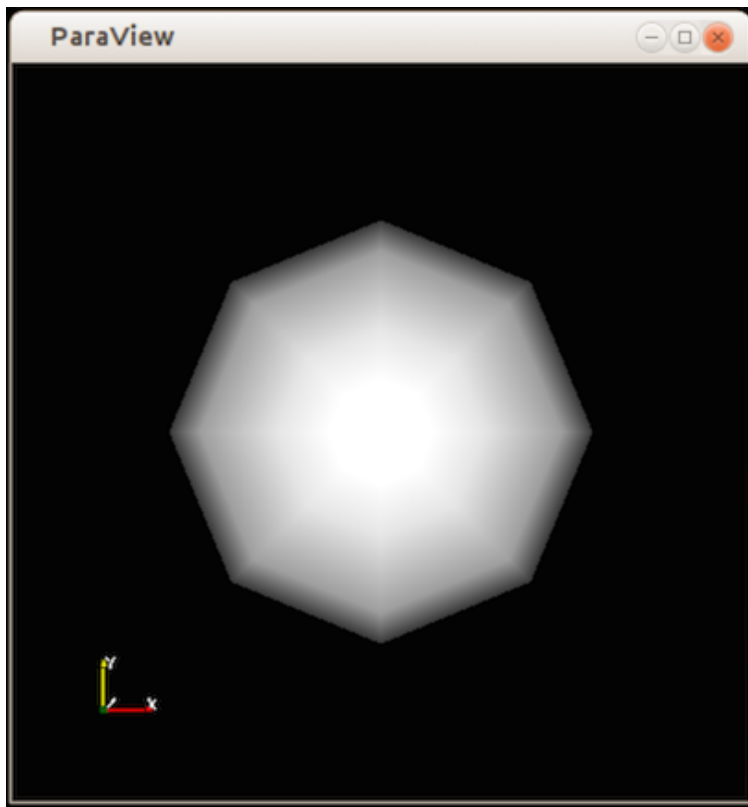


Fig. 1.5: Window showing result from the Python code.

Changing properties

To change the properties on the sphere source, you can use the `SetProperties` function.

```
# Set a single property on the active source.  
>>> SetProperties(Radius=1.0)  
  
# You can also set multiple properties.  
>>> SetProperties(Center=[1, 0, 0], StartTheta=100)
```

Similar to the `Properties` panel, `SetProperties` affects the active source. To query the current value of any property on the active source, use `GetProperty`.

```
>>> radius = GetProperty("Radius")  
>>> print(radius)  
1.0  
>>> center = GetProperty("Center")  
>>> print(center)  
[1.0, 0.0, 0.0]
```

`SetProperties` and `GetProperty` functions serve the same function as the `Properties` section of the `Properties` panel – they allow you to set and introspect the pipeline module properties for the active source. Likewise, for the `Display` section of the panel, or the display properties, we have the `SetDisplayProperties` and `GetDisplayProperty` functions.

```
>>> SetDisplayProperties(Opacity=0.5)  
>>> GetDisplayProperty("Opacity")  
0.5
```

Common Errors

Note how the property names for the `SetProperties` and `SetDisplayProperties` functions are not enclosed in double-quotes, while those for the `GetProperty` and `GetDisplayProperty` methods are.

In paraview, every time you hit `Apply` or change a display property, the UI automatically re-renders the view. In the scripting environment, you have to do this manually by calling the `Render` function every time you want to re-render and look at the updated result.

fixme{we're missing blurb about reset camera}.

Applying filters

Similar to creating a source, to apply a filter, you can simply create the filter by name.

```
# Create the 'Shrink' filter and connect it to the active source  
# which is the 'Sphere' instance.  
>>> Shrink()  
# As soon as the Shrink filter is created, it will now become the new active  
# source. All methods acting on active source now act on this filter instance  
# and not the Sphere instance created earlier.  
  
# Show the resulting data and render it.
```

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```
>>> Show()
>>> Render()
```

If you tried the above script, you'll notice the result isn't exactly what we expected. For some reason, the shrank cells are not visible. This is because we missed one stage: In paraview, the UI was smart enough to automatically hide the input dataset for the newly created filter after we hit apply. In the scripting interface, such operations are the user's responsibility. We should have hidden the sphere source from the view. We can use the Hide method, the counter part of Show , to hide the active source. But, now we have a problem – when we created the shrink filter, we changed the active source to be the shrink instance. Luckily, all the functions we discussed so far can take an optional first argument, which is the source or filter instance on which to operate. If provided, that instance is used instead of the active source. The solution is as follows:

```
# Get the input property for the active source, i.e. the input for the shrink.
>>> shrinksInput = GetProperty("Input")

# This is indeed the sphere instance we created earlier.
>>> print(shrinksInput)
<paraview.servermanager.Sphere object at 0x11d731e90>

# Hide the sphere instance explicitly.
>>> Hide(shrinksInput)

# Re-render the result.
>>> Render()
```

Alternatively, you could also get/set the active source using the GetActiveSource and SetActiveSource functions.

```
>>> shrinkInstance = GetActiveSource()
>>> print(shrinkInstance)
<paraview.servermanager.Shrink object at 0x11d731ed0>

# Get the input property for the active source, i.e. the input
# for the shrink.
>>> sphereInstance = GetProperty("Input")

# This is indeed the sphere instance we created earlier.
>>> print(sphereInstance)
<paraview.servermanager.Sphere object at 0x11d731e90>

# Change active source to sphere and hide it.
>>> SetActiveSource(sphereInstance)
>>> Hide()

# Now restore the active source back to the shrink instance.
>>> SetActiveSource(shrinkInstance)

# Re-render the result
>>> Render()
```

The result is shown in Fig. 1.6 .

SetActiveSource has same effect as changing the pipeline module, highlighted in the Pipeline Browser , by clicking on a different module.

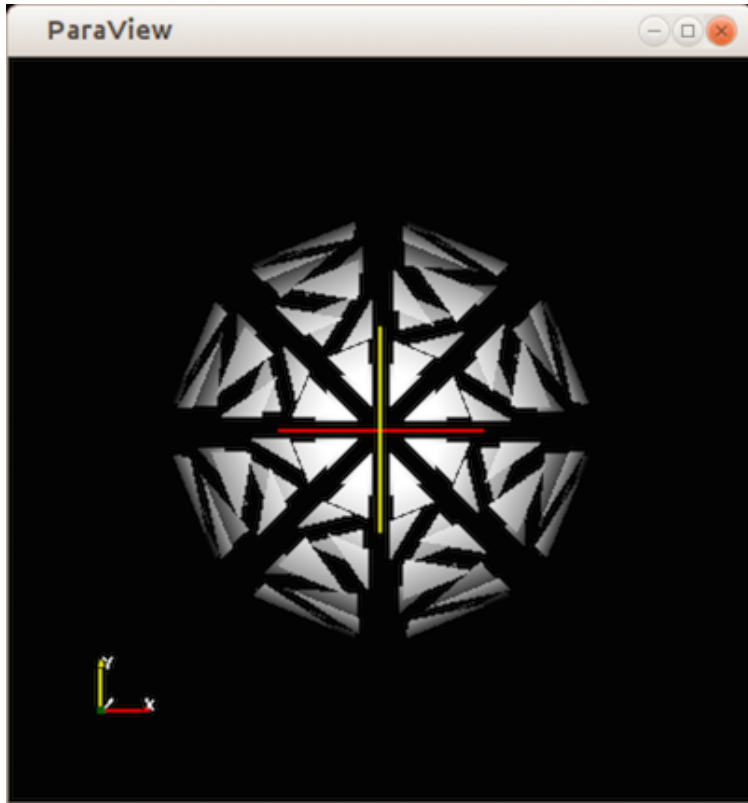


Fig. 1.6: Window showing result from the Python code after applying the shrink filter.

Alternative approach

Here's another way of doing something similar to what we did in the previous section for those familiar with Python and/or object-oriented programming. It's totally okay to stick with the previous approach.

```
>>> from paraview.simple import *
>>> sphereInstance = Sphere()
>>> sphereInstance.Radius = 1.0
>>> sphereInstance.Center[1] = 1.0
>>> print(sphereInstance.Center)
[0.0, 1.0, 0.0]

>>> sphereDisplay = Show(sphereInstance)
>>> view = Render()
>>> sphereDisplay.Opacity = 0.5

# Render function can take in an optional view argument, otherwise it
# will simply use the active view.
>>> Render(view)

>>> shrinkInstance = Shrink(Input=sphereInstance,
                           ShrinkFactor=1.0)
>>> print(shrinkInstance.ShrinkFactor)
1.0
>>> Hide(sphereInstance)
```

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```
>>> shrinkDisplay = Show(shrinkInstance)
>>> Render()
```

Updating the pipeline

When changing properties on the **Properties** panel in paraview, we noticed that the algorithm doesn't re-execute until you hit **Apply**. In reality, **Apply** isn't what's actually triggering the execution or the updating of the processing pipeline. What happens is that **Apply** updates the parameters on the pipeline module and causes the view to render. If the output of the pipeline module is visible in the view, or if the output of any filter connected to it downstream is visible in the view, **ParaView** will determine that the data rendered is obsolete and request the pipeline to re-execute. It implies that if that pipeline module (or any of the filters downstream from it) is not visible in the view, **ParaView** will have no reason to re-execute the pipeline, and the pipeline module will not be updated. If, later on, you do make this module visible in the view, **ParaView** will automatically update and execute the pipeline. This is often referred to as *demand-driven pipeline execution*. It makes it possible to avoid unnecessary module executions.

In paraview, you can get by without ever noticing this since the application manages pipeline updates automatically. In pypython too, if your scripts are producing renderings in views, you'd never notice this as long as you remember to call **Render**. However, you may want to write scripts to produce transformed datasets or to determine data characteristics. In such cases, since you may never create a view, you'll never be seeing the pipeline update, no matter how many times you change the properties.

Accordingly, you must use the **UpdatePipeline** function. **UpdatePipeline** updates the pipeline connected to the active source (or only until the active source, i.e., anything downstream from it, won't be updated).

```
>>> from paraview.simple import *
>>> sphere = Sphere()

# Print the bounds for the data produced by sphere.
>>> print(sphere.GetDataInformation().GetBounds())
(1e+299, -1e+299, 1e+299, -1e+299, 1e+299, -1e+299)
# The bounds are invalid -- no data has been produced yet.

# Update the pipeline explicitly on the active source.
>>> UpdatePipeline()

# Alternative way of doing the same but specifying the source
# to update explicitly.
>>> UpdatePipeline(proxy=sphere)

# Let's check the bounds again.
>>> sphere.GetDataInformation().GetBounds()
(-0.48746395111083984, 0.48746395111083984, -0.48746395111083984, 0.48746395111083984, -0.5, 0.5)

# If we call UpdatePipeline() again, this will have no effect since
# the pipeline hasn't been modified, so there's no need to re-execute.
>>> UpdatePipeline()
>>> sphere.GetDataInformation().GetBounds()
(-0.48746395111083984, 0.48746395111083984, -0.48746395111083984, 0.48746395111083984, -0.5, 0.5)

# Now let's change a property.
```

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```
>>> sphere.Radius = 10

# The bounds won't change since the pipeline hasn't re-executed.
>>> sphere.GetDataInformation().GetBounds()
(-0.48746395111083984, 0.48746395111083984, -0.48746395111083984, -
 0.5, 0.5)

# Let's update and see:
>>> UpdatePipeline()
>>> sphere.GetDataInformation().GetBounds()
(-9.749279022216797, 9.749279022216797, -9.749279022216797, 9.749279022216797, -10.0, 10.
 0)
```

We will look at the `sphere.GetDataInformation` API in [Section 1.3.3](#) in more detail.

For temporal datasets, `UpdatePipeline` takes in a time argument, which is the time for which the pipeline must be updated.

```
# To update to time 10.0:
>>> UpdatePipeline(10.0)

# Alternative way of doing the same:
>>> UpdatePipeline(time=10.0)

# If not using the active source:
>>> UpdatePipeline(10.0, source)
>>> UpdatePipeline(time=10.0, proxy=source)
```

1.1.6 Scripting in paraview

The Python Shell

The paraview application also provides access to an internal shell, in which you can enter Python commands and scripts exactly as with `pypython`. To access the Python shell in the GUI, use the *View > Python Shell* menu option. A dialog will pop up with a prompt exactly like `pypython`. You can try inputting commands from the earlier section into this shell. As you type each of the commands, you will see the user interface update after each command, e.g., when you create the `sphere` source instance, it will be shown in the `Pipeline Browser`. If you change the active source, the `Pipeline Browser` and other UI components will update to reflect the change. If you change any properties or display properties, the `Properties` panel will update to reflect the change as well!

Did You Know?

The `Python Shell` in `paraview` supports auto-completion for functions and instance methods. Try hitting the Tab key after partially typing any command (as shown in [Fig. 1.7](#)).

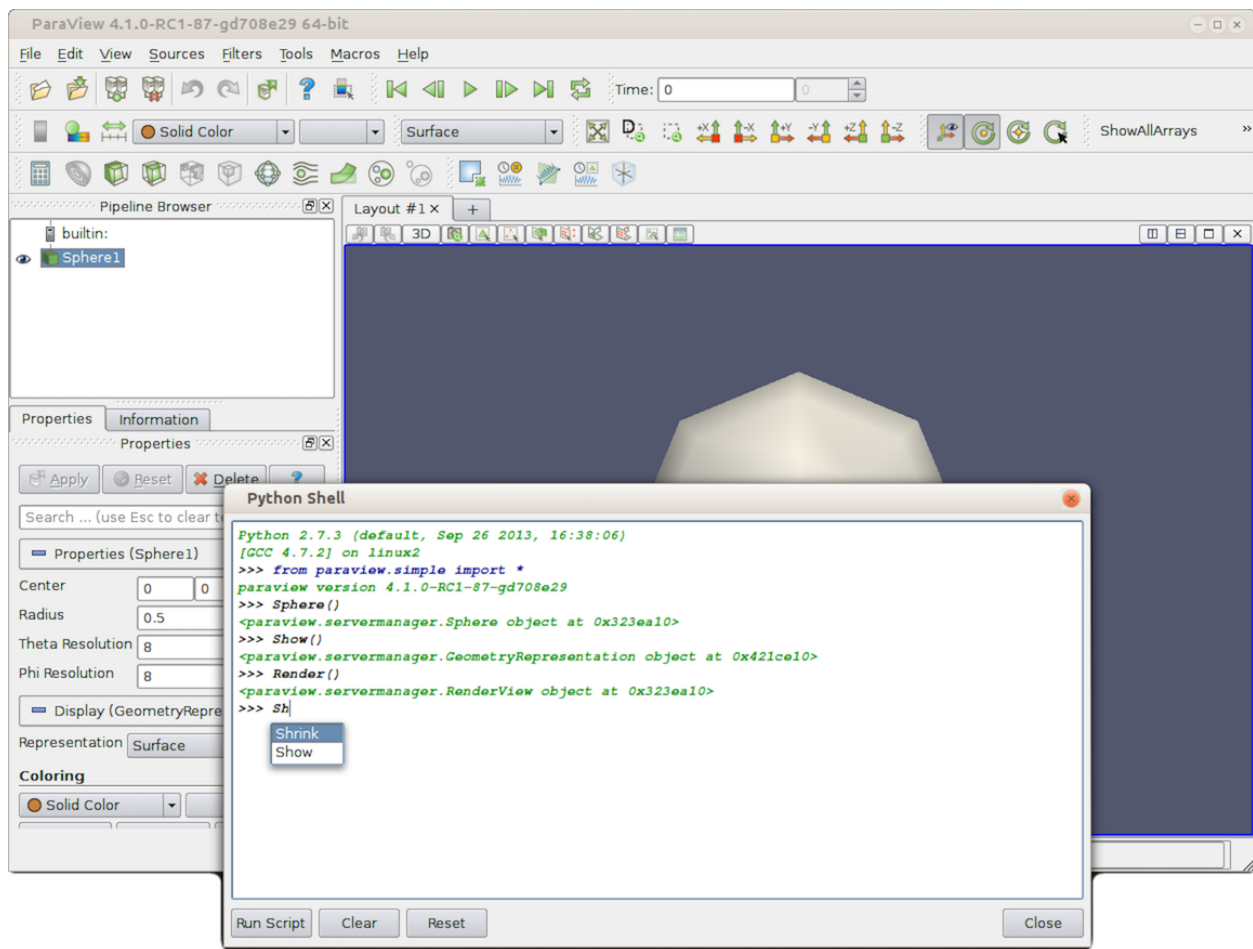


Fig. 1.7: Python Shell in paraview provides access to the scripting.

Tracing actions for scripting

This guide provides a fair overview of **ParaView**'s Python API. However, there will be cases when you just want to know how to complete a particular action or sequence of actions that you can do with the GUI using a Python script instead. To accomplish this, `paraview` supports tracing your actions in the UI as a Python script. Simply start tracing by clicking on *Tools > Start Trace*. `paraview` now enters a mode where all your actions (or at least those relevant for scripting) are monitored. Any time you create a source or filter, open data files, change properties and hit **Apply**, interact with the 3D scene, or save screenshots, etc., your actions will be monitored. Once you are done with the series of actions that you want to script, click *Tools > Stop Trace*. `paraview` will then pop up an editor window with the generated trace. This will be the Python script equivalent for the actions you performed. You can now save this as a script to use for batch processing.

1.2 Loading Data

In a visualization pipeline, data sources bring data into the system for processing and visualization. Sources, such as the **Sphere** source (accessible from the **Sources** menu in `paraview`), programmatically create datasets for processing. Another type of data sources are readers. Readers can read data written out in disk files or other databases and bring it into **ParaView** for processing. **ParaView** includes readers that can read several of the commonly used scientific data formats. It's also possible to write plugins that add support for new or proprietary file formats.

ParaView provides several sample datasets for you to get started. You can download an archive with several types of data files from the download page at <https://www.paraview.org/download> under the **Data** section.

1.2.1 Opening data files in paraview

To open a data file in `paraview`, you use the **Open File** dialog. This dialog can be accessed from the **File > Open** menu or by using the  button in the **Main Controls** toolbar. You can also use the keyboard shortcut **CTRL + O** (or **+ O**) to open this dialog.

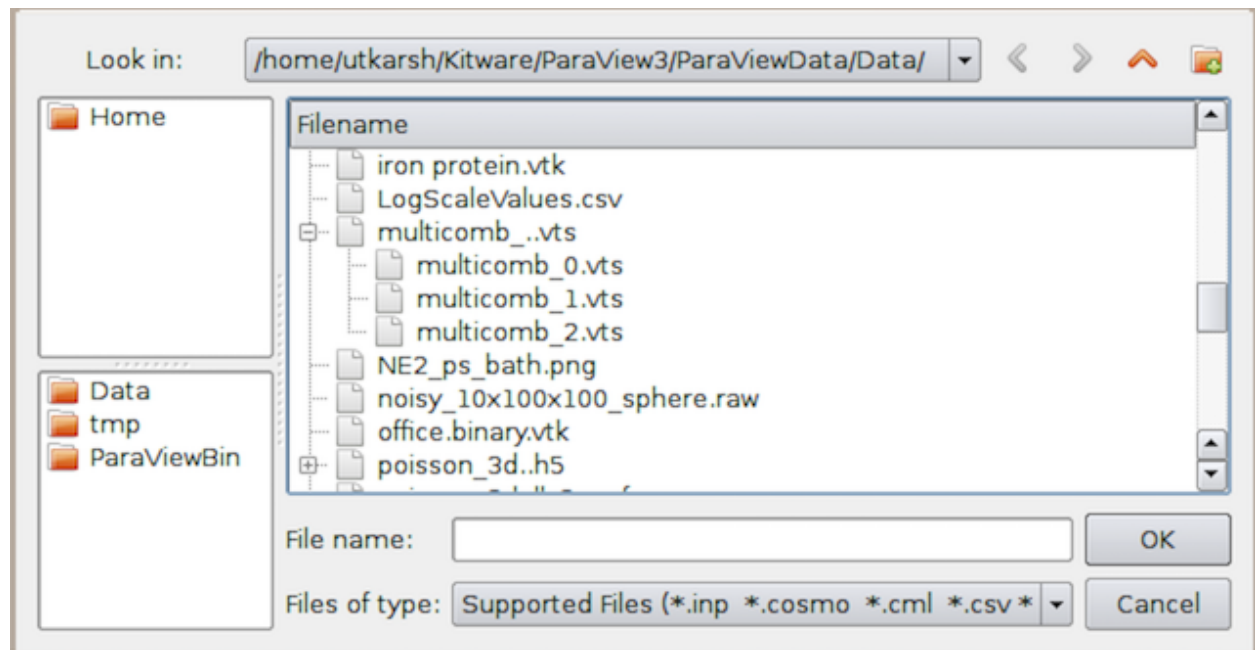
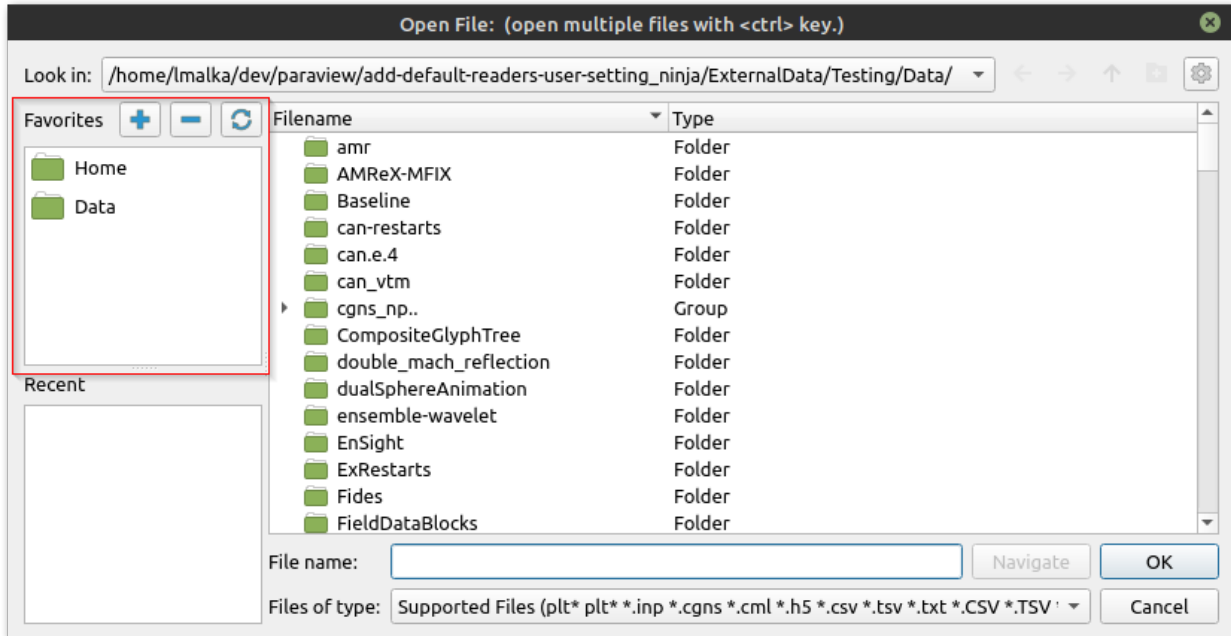


Fig. 1.8: **Open File** dialog in `paraview` for opening data (and other) files.

The **Open File** dialog allows you to browse the file system on the data processing nodes. This will become clear when we look at using **ParaView** for remote visualization. While several of the UI elements in this dialog are obvious such as navigating up the current directory, creating a new directory, and navigating back and forth between directories, which can all be done with the standard system shortcuts like **CTRL + N** (or **+** **N**) to create a directory or **Alt + ↑** to go to the parent directory, there are a few things to note.



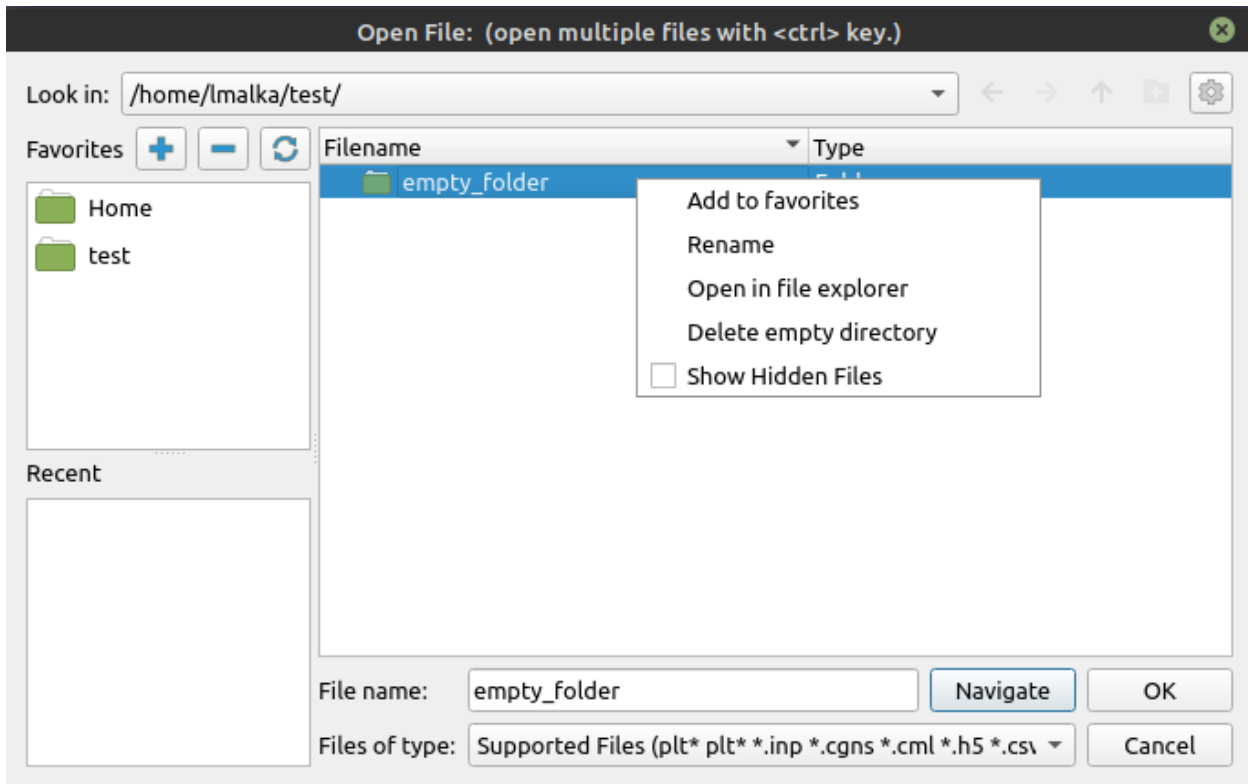
- The **Favorites** pane shows some platform-specific common locations such as the home directory and desktop. The favorites directories can be customized from the buttons above it, which respectively adds the current directory to the favorites, removes the current directory from the favorites, and resets all the favorites to the system default. Another way to change which directories are in the favorites is to use the right-click context menus. Right-clicking any directory in the main list will display a menu with the option to add it to the favorites. A right-click on a favorite in the favorites list displays the option to remove it.
- The **Recent Directories** pane shows a few of the most recently used directories.

You can browse to the directory containing your datasets and either select the file and hit **Ok** or simply double click on the file to open it. You can also select multiple files using the **CTRL** (or **+**) key. This will open each of the selected files separately.

Right-clicking the files list will display a few options, depending on what was right-clicked.

- The **Show Hidden Files** is always visible, and can be checked to display the hidden files and directories.
- The **Open in file explorer** is also always present and either opens in the system file explorer the selected directory if one was right-clicked, or opens the current directory if a file or nothing was selected.
- Selecting a file or a directory adds the **Rename** option.
- Selecting a folder adds the **Add to favorites** option which adds the selected directory to the **Favorites** pane on the left.
- Selecting an empty directory provides the **Delete empty directory** option.

When a file is opened, **paraview** will create a reader instance of the type suitable for the selected file based on its extension. The reader will simply be another pipeline module, similar to the source we created in [Section 1.1](#). From this point forward, the workflow will be the same as we discussed in [Section 1.1.4](#) : You adjust the reader properties, if needed, and hit **Apply**. **paraview** will then read the data from the file and render it in the view.



If you selected multiple files using the CTRL (or) key, paraview will create multiple reader modules. When you hit Apply , all of the readers will be executed, and their data will be shown in the view.

Did you know?

This ability to hit the Apply button once to accept changes on multiple readers applies to other pipeline modules, including sources and filters. In general, you can change properties for multiple modules in a pipeline, and hit Apply to accept all of the changes at once. It is possible to override this behavior from the Settings dialog.

Reader selection depending on the selected file types

When selecting files, the current category for the “Files of type” field (Fig. 1.10) changes the way the reader will be chosen.

- If the “Supported Files” value is selected, if only one reader is available for the type of file, it will be selected automatically. If multiple readers can be used, the “Open Data With...” dialog (Fig. 1.11) will appear to choose which reader to use. The “Set reader as default” button can be clicked to automatically use this reader when one of its pattern matches a file that could otherwise be opened by multiple readers. When a reader is set as default, it will be used automatically for files that match its patterns.
- If the “All Files” value is selected, the same dialog will be displayed with all the existing readers. If you picked an incorrect reader, however, you’ll get error messages either when the reader module is instantiated or after you hit Apply. In either case, you can simply Delete the reader module and try opening the file again, this time choosing a different reader. When clicking “Set reader as default”, a line edit will be displayed to set the custom pattern to use for this reader (Fig. 1.12). This is a standard wildcard pattern, and multiple patterns can be used by separating them with spaces.

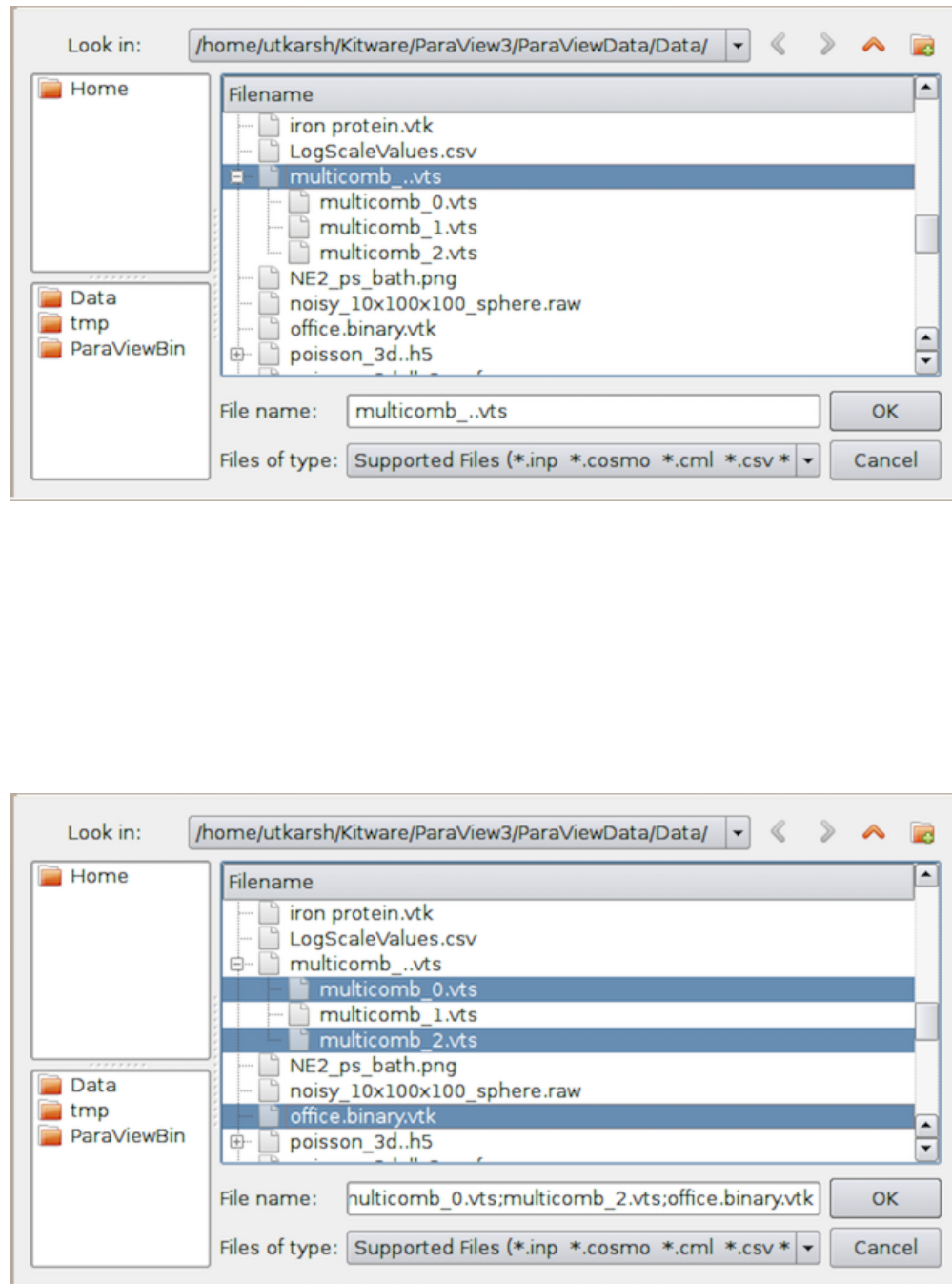


Fig. 1.9: The Open File dialog can be used to select a temporal file series (top) or select multiple files to open separately (bottom).

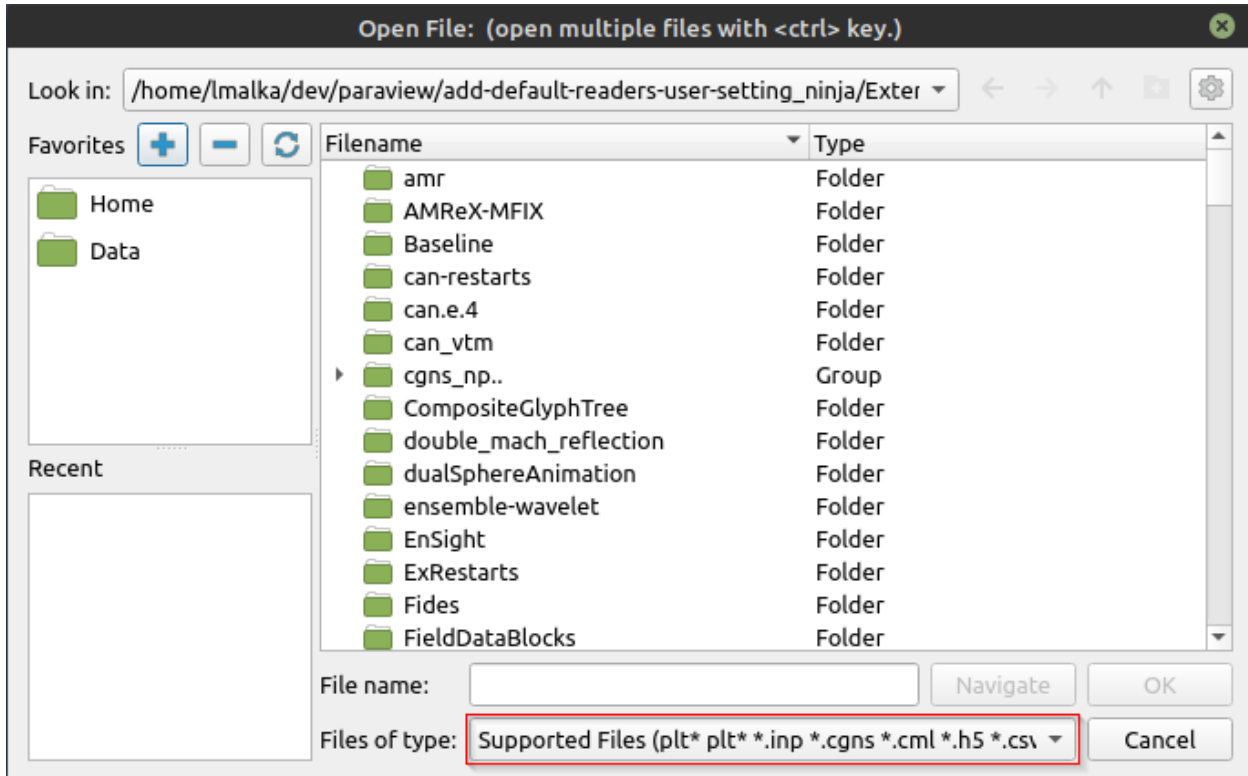


Fig. 1.10: File dialog “Files of type” to filter which files to show and to change how the reader will be chosen.

- If a specific reader is selected, for example “PNG Image Files”, this reader will be always be used automatically, even if other readers would be available or if the file is matching a pattern present in the “Default reader details” settings.

Error messages in paraview are shown in the `Output Messages` window (Fig. 1.13). It is accessible from the `View > Output Messages` menu. Whenever there’s a new error message, paraview will automatically pop open this window and raise it to the top. This window can be attached, or docked, in the main window so that it is visible with the other user interface elements without covering them up.

The default readers settings can be seen and modified in the `Edit -> Settings` menu, and `Miscellaneous` tab (Fig. 1.14).

Handling temporal file series

Most datasets produced by scientific simulation runs are temporal in nature. File formats differ in how this information is saved in the file. While several file formats support saving multiple timesteps in the same file, others save them out as a sequence of files, known as a file series

The Open File dialog automatically detects file series and shows them as a grouped element, as shown in Fig. 1.9. To load the file series, simply select the group, and hit Ok . You can also open a single file in the series as a regular file. To do so, open the file group and select the file you want to open.

paraview automatically detects several of the commonly-known file naming patterns used for indicating a file series. These include:

fooN.vtk	fooN.vtk	Nfoo.vtk	foo.vtk.N
foo_N.vtk	foo.N.vtk	N.foo.vtk	foo.vtksN

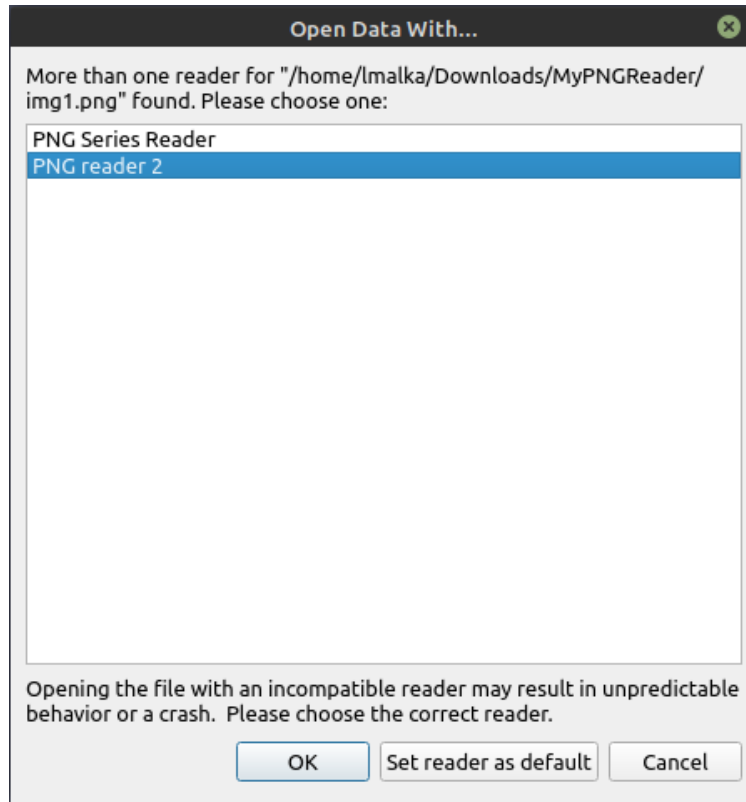


Fig. 1.11: Open Data With... dialog shown to manually choose the reader to use for a file with multiple available readers.

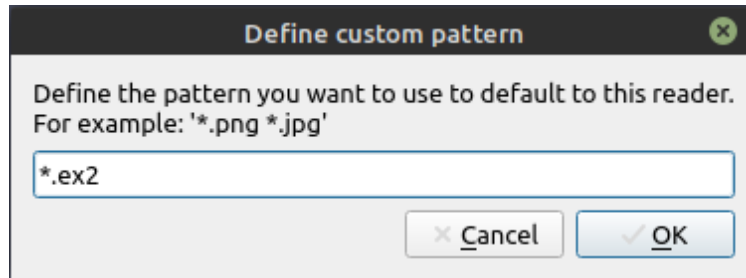


Fig. 1.12: The input to define the pattern to add to the settings to use this reader by default.

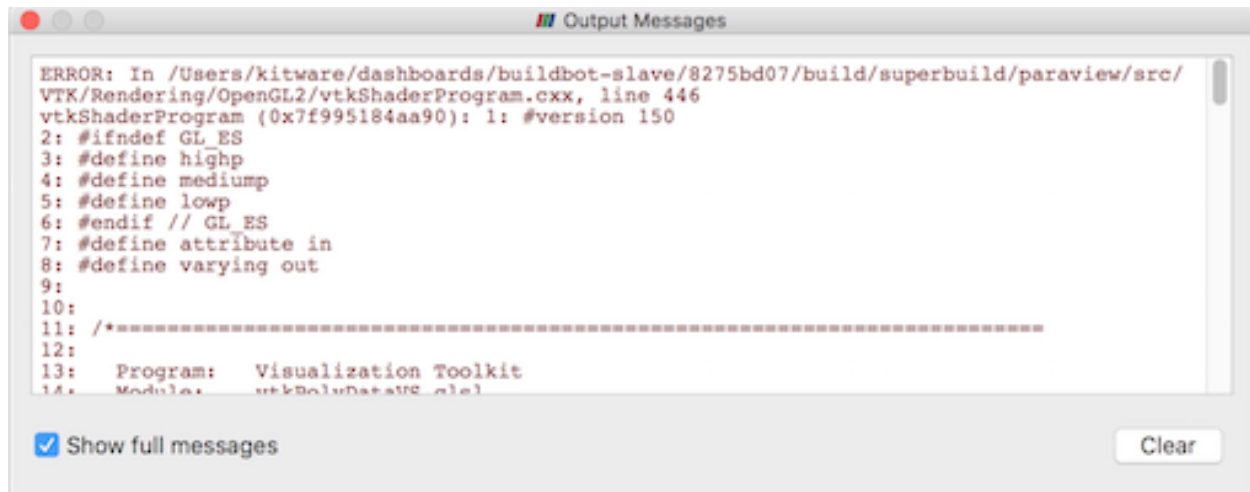


Fig. 1.13: The Output Messages window is used to show errors, warnings, and other messages raised by the application.

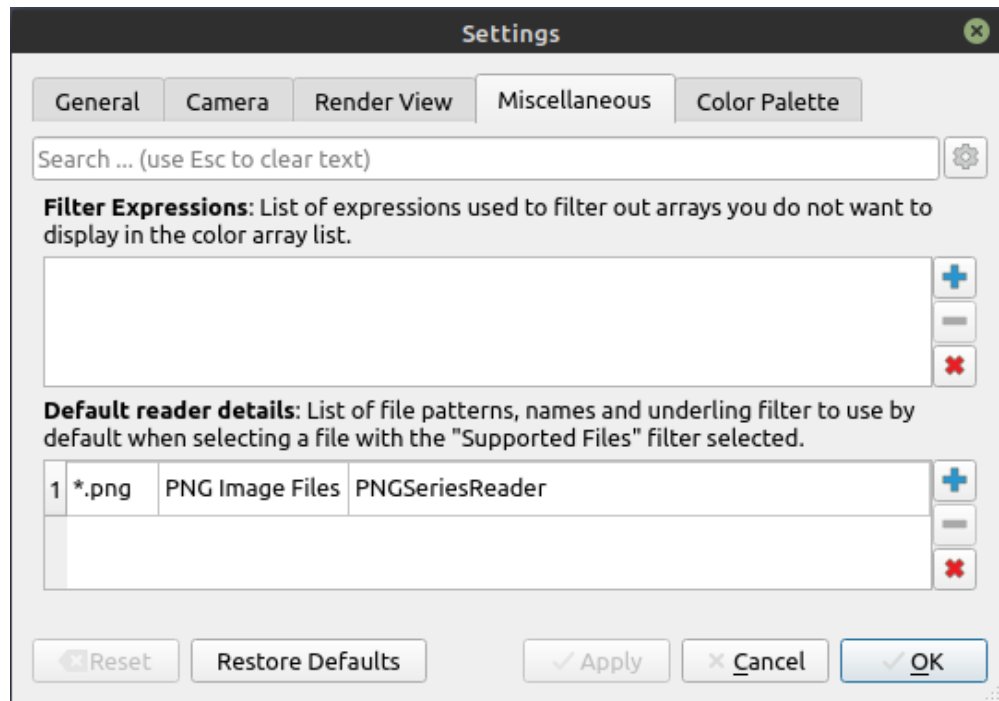


Fig. 1.14: Default reader settings.

where *foo* could be any filename, *N* is a numeral sequence (with any number of leading zeros), and *vtk* could be any extension.

Dealing with time

When you open a dataset with time, either as a file series or in a file format that natively supports time, `paraview` will automatically setup an animation for you so that you can play through each of the time steps in the dataset by using the

▶ button on the VCR Controls toolbar (Fig. 1.15). You can change or modify this animation and further customize it, as discussed in Chapter [Section 1.7](#).



Fig. 1.15: VCR Controls toolbar for interacting with an animation.

Reopening previously opened files

`paraview` remembers most recently opened files (or file series). Simply use the *File > Recent Files* menu. `paraview` also remembers the reader type selected for files with unknown extensions or for occasions when multiple reader choices were available.

Opening files using command line options

`paraview` provides a command line option that can be used to open datasets on startup.

```
> paraview --data=.../ParaViewData/Data/can.ex2
```

This is equivalent to opening a `can.ex2` data file from the *Open File* dialog. The same set of follow-up actions happen. For example, `paraview` will try to locate a reader for the file, create that reader, and wait for you to hit *Apply*.

To open a file series, simply replace the numbers in the file name sequence by a `.`. For example, to open a file series named `my0.vtk`, `my1.vtk` ... `myN.vtk`, use `my..vtk`.

```
> paraview --data=.../ParaViewData/Data/my..vtk
```

Common properties for readers

ParaView uses different reader implementations for different file formats. Each of these have different properties available to you to customize how the data is read and can vary greatly depending on the capabilities of the file format itself or the particular reader implementation. Let's look at some of the properties commonly available in readers.

Selecting data arrays

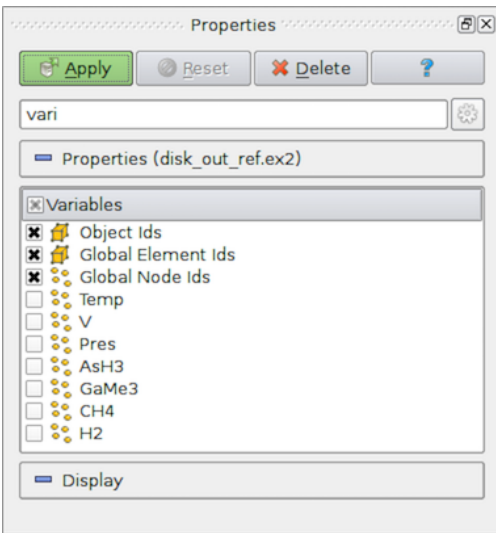
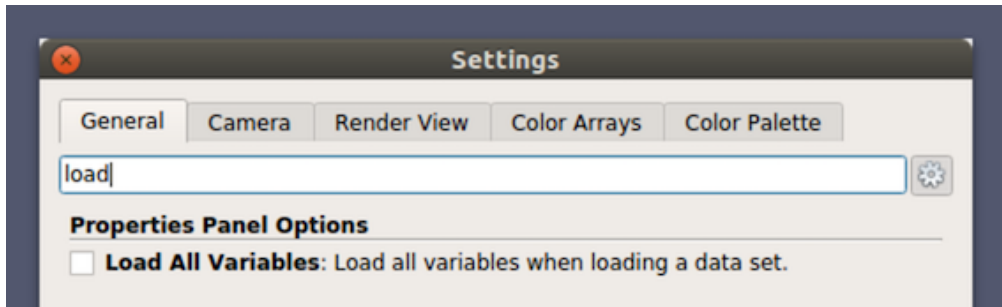


Fig. 1.16: Array selection widget for selecting array to load from a data file.

One of the most common properties on readers is one that allows you to select the data arrays (cell centered, point centered, or otherwise, if applicable) to be loaded. Often times, loading only the data arrays you know you are going to use in the visualization will save memory, as well as processing time, since the reader does not have to read in those data arrays, and the filters in the pipeline do not have to process them.

Did you know?

You can change paraview's default behavior to load all available data arrays by selecting the **Load All Variables** checkbox under **Settings/Properties Panel Options/Advanced**.



The user interface for selecting the arrays to load is simply a list with the names of the arrays and a checkbox indicating whether that array is to be loaded or not (Fig. 1.16). Icons, such as and , are often used in this widget to give you an indication of whether the array is cell-centered or point-centered, respectively.

If you initially de-select an array, but then as you're setting up your visualization pipeline realize that you need that data array, you can always go back to the Properties page for the reader by making the reader active in the Pipeline Browser and then changing the array selection. **ParaView** will automatically re-execute any processing pipeline set up on the reader with this new data array.

Common Errors

Remember to hit **Apply** (or use **Auto Apply**) after changing the array selection for the change to take effect.

Sometimes the list of data arrays can get quite large, and it can become cumbersome to find the array for which you are looking. To help with such situations, paraview provides a mechanism to search lists. Click inside the widget to make it get the *focus*. Then type **CTRL + F** (or **+ F**) to get a search widget. Now you can type in the text to search. Matching rows will be highlighted (Fig. 1.17).

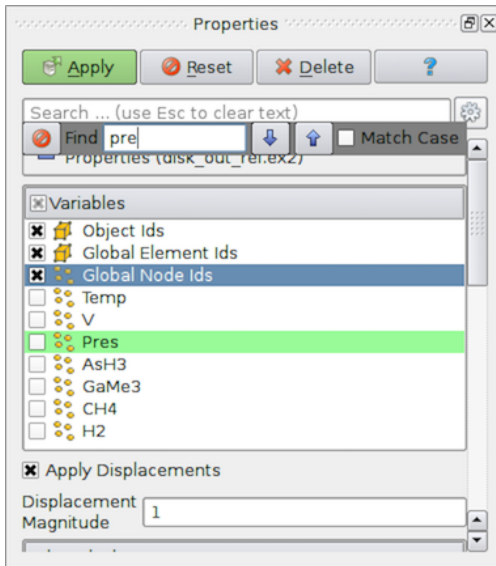


Fig. 1.17: To search through large lists in paraview, you can use **CTRL + F**.

Did you know?

The ability to search for items in an array selection widget also applies to other list and tree widgets in the paraview UI. Whenever you see a widget with a large number of entries in a list, table, or tree fashion, try using **CTRL + F** (or **+ F**).

1.2.2 Opening data files in pypython

To open data files using the scripting interface, **ParaView** provides the `OpenDataFile` function.

```
>>> reader = OpenDataFile("../ParaViewData/Data/can.ex2")
>>> if reader:
...     print("Success")
... else:
...     print("Failed")
... 
```

`OpenDataFile` will try to determine an appropriate reader based on the file extension, just like **paraview**. If no reader is determined, `None` is returned. If multiple readers can open the file, however, `OpenDataFile` simply picks the first reader. If you explicitly want to create a specific reader, you can always create the reader by its name, similar to other sources and filters.

```
>>> reader = ExodusIIReader(FileName=".../ParaViewData/Data/can.ex2")
```

To find out information about the reader created and the properties available on it, you can use the `help` function.

```
>>> reader = ExodusIIReader(FileName=".../ParaViewData/Data/can.ex2")
```

```
>>> help(reader)
```

Help on ExodusIIReader in module paraview.servermanager object:

```
class ExodusIIReader(ExodusIIReaderProxy)
| The Exodus reader loads
| Exodus II files and produces an unstructured grid output.
| The default file extensions are .g, .e, .ex2, .ex2v2,
| .exo, .gen, .exoII, .exii, .0, .00, .000, and .0000. The
| file format is described fully at:
| http://endo.sandia.gov/SEACAS/Documentation/exodusII.pdf.
| ...
|
```

```
|-----
```

| Data descriptors defined here:

```
| AnimateVibrations
```

| If this flag is on and HasModeShapes is also on, then
| this reader will report a continuous time range [0,1] and
| animate the displacements in a periodic sinusoid. If this
| flag is off and HasModeShapes is on, this reader ignores
| time. This flag has no effect if HasModeShapes is off.

```
| ApplyDisplacements
```

| Geometric locations can include displacements. When this
| option is on, the nodal positions are 'displaced' by the
| standard exodus displacement vector. If displacements are
| turned 'off', the user can explicitly add them by applying
| a warp filter.

```
| ...
|
```

Did you know?

The `help` function can be used to get information about properties available on any source or filter instance. It not only lists the properties, but also provides information about how they affect the pipeline module. `help` can also be used on functions. For example:

```
>>> help(OpenDataFile)
```

Help on function OpenDataFile in module paraview.simple:

```
OpenDataFile(filename, \*\*extraArgs)
```

Creates a reader to read the given file, if possible.
This uses extension matching to determine the best reader
possible. If a reader cannot be identified, then this
returns None.

Handling temporal file series

Unlike paraview, pvython does not automatically detect and load file series . You have to explicitly list the filenames in the series and pass that to the OpenDataFile call.

```
# Create a list with the names of all the files in the file series in
# correct order.
>>> files = [".../Data/multicomb_0.vts",
             ".../Data/multicomb_1.vts",
             ".../Data/multicomb_2.vts"]
>>> reader = OpenDataFile(files)
```

Dealing with time

Similar to paraview, if you open a time series or a file with multiple timesteps, pvython will automatically set up an animation for you to play through the timesteps.

```
>>> files = [".../Data/multicomb_0.vts",
             ".../Data/multicomb_1.vts",
             ".../Data/multicomb_2.vts"]
>>> reader = OpenDataFile(files)
>>> Show()
>>> Render()

# Get access to the animation scene.
>>> scene = GetAnimationScene()
# Now you use the API on the scene when doing things such as playing
# the animation, stepping through it, etc.

# This will simply play through the animation once and stop. Watch
# the rendered view after you hit 'Enter.'
>>> scene.Play()
```

Common properties on readers

Selecting data arrays

For those properties on readers that allow you to control what to read in from the file such as point data arrays, cell data arrays, or data blocks, paraview uses a selection widget, as seen in [Section 1.2.1](#). Likewise, pvython provides an API that allows you to determine the available options and then select/deselect them.

The name of the property that allows you to make such selections depends on the reader itself. When in doubt, use the tracing capabilities in paraview ([Section 1.1.6](#)) to figure it out. You can also use help ([Section 1.2.2](#)).

ExodusIIReader has a PointVariables property that can be used to select the point data arrays to load. Let's use this as an example.

```
# Open an ExodusII data file.
>>> reader = OpenDataFile(".../Data/can.ex2")

# Alternatively, you can explicitly create the reader instance as:
>>> reader = ExodusIIReader(FileName = ".../Data/can.ex2")
```

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```

# To query/print the current status for `PointVariables` property,
# we do what we would have done for any other property:
>>> printGetProperty("PointVariables"))
['DISPL', 'VEL', 'ACCL']

# An alternative way of doing the same is as follows:
>>> print(reader.PointVariables)
['DISPL', 'VEL', 'ACCL']

# To set the property, simply set it to list containing the names to
# enable, e.g., if we want to read only the 'DISPL' array, we do
# the following:
>>> SetProperties(PointVariables=['DISPL'])

# Or using the alternative way for doing the same:
>>> reader.PointVariables = ['DISPL']

# Now, the new value for PointVariables is:
>>> print(reader.PointVariables)
['DISPL']

# To determine the array available, use:
>>> print(reader.PointVariables.Available)
['DISPL', 'VEL', 'ACCL']
# These are the arrays available in the file.

```

Changing `PointVariables` only changes the value on the property. The reader does not re-execute until a re-execution is requested either by calling `Render` or by explicitly updating the pipeline using `UpdatePipeline`.

```

>>> reader.PointVariables = ['DISPL', 'VEL', 'ACCL']

# Assuming that the reader is indeed the active source, let's update
# the pipeline:
>>> UpdatePipeline()

# Or you can use the following form if you're unsure of the active
# source or just do not want to worry about it.
>>> UpdatePipeline(proxy=reader)

# Print the list of point arrays read in.
>>> print(reader.PointData[:])
[Array: ACCL, Array: DISPL, Array: GlobalNodeId, Array: PedigreeNodeId, Array: VEL]

# Change the selection.
>>> reader.PointVariables = ['DISPL']

# Print the list of point arrays read in, nothing changes!
>>> print(reader.PointData[:])
[Array: ACCL, Array: DISPL, Array: GlobalNodeId, Array: PedigreeNodeId, Array: VEL]

# Update the pipeline.

```

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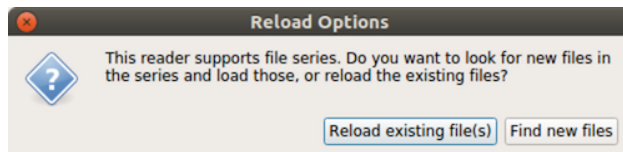
```
>>> UpdatePipeline()

# Now the arrays read in has indeed changed as we expected.
>>> print(reader.PointData[:])
[Array: DISPL, Array: GlobalNodeId, Array: PedigreeNodeId]
```

We will cover the `reader.PointData` API in more details in [Section 1.3.3](#).

1.2.3 Reloading files

While ParaView is often used after the simulation has generated all the data, it is not uncommon to use ParaView to inspect data files as they are being written out by the simulation. In such cases, the simulation may either be modifying existing file(s) with new timesteps or creating new files for each timestep. In such cases, you may want to *refresh* ParaView to make it aware of the changes. In paraview, this can be done using **Reload Files**. When the reader is active, you can use the *File > Reload Files* menu to request the reader to refresh. paraview will prompt you to choose whether to reload the existing file(s) or look for new files in the file series, as shown in [Fig. 1.18](#). Click on `Reload existing file(s)`, to force the reader to re-read the files already opened. This is useful in cases where the simulation may have modified existing file(s). Use `Find new files` to make the reader aware of any new files in the file series.



[Fig. 1.18](#): The `Reload Options` dialog allows you to choose how to refresh the reader.

Similar to `paraview`, in `pypython`, you use `ReloadFiles` to reload existing files, and `ExtendFilesSeries` to look for new files in a file series.

```
# For file being modified in place per timestep
>>> reader = OpenDataFile(file)
...
>>> ReloadFiles(reader)

# For files being generated per timestep
>>> reader = OpenDataFile(file)
...
>>> ExtendFilesSeries(reader)
```

1.3 Understanding Data

1.3.1 VTK data model

To use **ParaView** effectively, you need to understand the **ParaView** data model. **ParaView** uses VTK, the Visualization Toolkit, to provide the visualization and data processing model. This chapter briefly introduces the VTK data model used by **ParaView**. For more details, refer to one of the VTK books.

The most fundamental data structure in VTK is a data object. Data objects can either be scientific datasets, such as rectilinear grids or finite elements meshes (see below), or more abstract data structures, such as graphs or trees. These datasets are formed from smaller building blocks: mesh (topology and geometry) and attributes.

Mesh

Even though the actual data structure used to store the mesh in memory depends on the type of the dataset, some abstractions are common to all types. In general, a mesh consists of vertices (points) and cells (elements, zones). Cells are used to discretize a region and can have various types such as tetrahedra, hexahedra, etc. Each cell contains a set of vertices. The mapping from cells to vertices is called the connectivity. Note that even though it is possible to define data elements such as faces and edges, VTK does not represent these explicitly. Rather, they are implied by a cell's type and by its connectivity. One exception to this rule is the arbitrary polyhedron, which explicitly stores its faces. [Fig. 1.19](#) is an example mesh that consists of two cells. The first cell is defined by vertices (0, 1, 3, 4), and the second cell is defined by vertices (1, 2, 4, 5). These cells are neighbors because they share the edge defined by the points (1, 4).

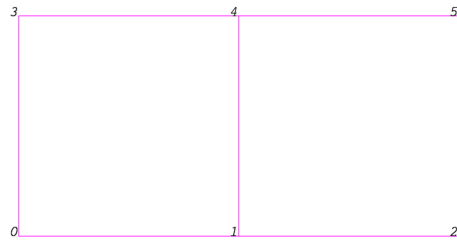


Fig. 1.19: Example of a mesh.

A mesh is fully defined by its topology and the spatial coordinates of its vertices. In VTK, the point coordinates may be implicit, or they may be explicitly defined by a data array of dimensions (*number_of_points* × 3).

Attributes (fields, arrays)

An attribute (or a data array or field) defines the discrete values of a field over the mesh. Examples of attributes include pressure, temperature, velocity, and stress tensor. Note that VTK does not specifically define different types of attributes. All attributes are stored as data arrays, which can have an arbitrary number of components. **ParaView** makes some assumptions in regards to the number of components. For example, a 3-component array is assumed to be an array of vectors. Attributes can be associated with points or cells. It is also possible to have attributes that are not associated with either. [Fig. 1.20](#) demonstrates the use of a point-centered attribute. Note that the attribute is only defined on the vertices. Interpolation is used to obtain the values everywhere else. The interpolation functions used depend on the cell type. See the VTK documentation for details.

[Fig. 1.21](#) demonstrates the use of a cell-centered attribute. Note that cell-centered attributes are assumed to be constant over each cell. Due to this property, many filters in VTK cannot be directly applied to cell-centered attributes. It is normally required to apply a **Cell Data to Point Data** filter. In **ParaView**, this filter is applied automatically, when necessary.

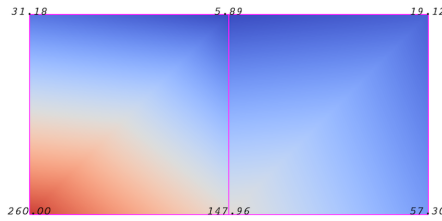


Fig. 1.20: Point-centered attribute in a data array or field.

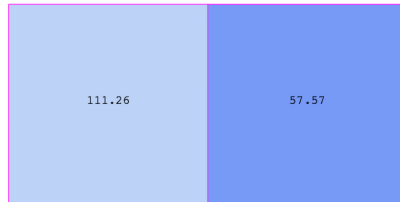


Fig. 1.21: Cell-centered attribute.

Uniform rectilinear grid (image data)

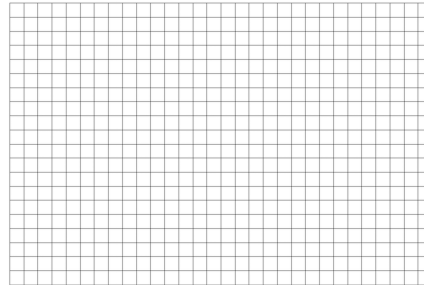


Fig. 1.22: Example uniform rectilinear grid.

A uniform rectilinear grid, or image data, defines its topology and point coordinates implicitly (Fig. 1.22). To fully define the mesh for an image data, VTK uses the following:

- *Extents* - These define the minimum and maximum indices in each direction. For example, an image data of extents (0, 9), (0, 19), (0, 29) has 10 points in the x-direction, 20 points in the y-direction, and 30 points in the z-direction. The total number of points is $10 \times 20 \times 30$.
- *Origin* - This is the position of a point defined with indices (0, 0, 0).
- *Spacing* - This is the distance between each point. Spacing for each direction can be defined independently.

The coordinate of each point is defined as follows: $coordinate = origin + index \times spacing$ where $coordinate$, $origin$, $index$, and $spacing$ are vectors of length 3.

Note that the generic VTK interface for all datasets uses a flat index. The (i, j, k) index can be converted to this flat index as follows: $idx_flat = k \times (npts_x \times npts_y) + j \times nptr_x + i$.

A uniform rectilinear grid consists of cells of the same type. This type is determined by the dimensionality of the dataset (based on the extents) and can either be vertex (0D), line (1D), pixel (2D), or voxel (3D).

Due to its regular nature, image data requires less storage than other datasets. Furthermore, many algorithms in VTK have been optimized to take advantage of this property and are more efficient for image data.

Rectilinear grid

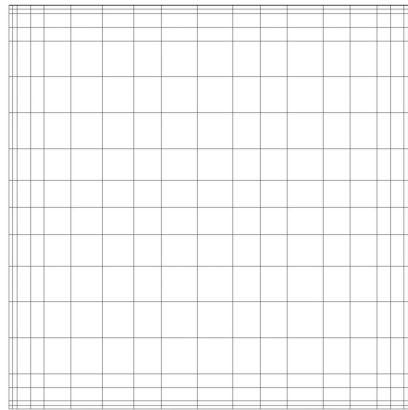


Fig. 1.23: Rectilinear grid.

A rectilinear grid, such as Fig. 1.23, defines its topology implicitly and point coordinates semi-implicitly. To fully define the mesh for a rectilinear grid, VTK uses the following:

- *Extents* - These define the minimum and maximum indices in each direction. For example, a rectilinear grid of extents $(0, 9)$, $(0, 19)$, $(0, 29)$ has 10 points in the x-direction, 20 points in the y-direction, and 30 points in the z-direction. The total number of points is $10 \times 20 \times 30$.
- *Three arrays defining coordinates in the x-, y- and z-directions* - These arrays are of length $npts_x$, $npts_y$, and $npts_z$. This is a significant savings in memory, as the total memory used by these arrays is $npts_x + npts_y + npts_z$ rather than $npts_x \times npts_y \times npts_z$.

The coordinate of each point is defined as follows:

$$coordinate = (coordinate_array_x(i), coordinate_array_y(j), coordinate_array_z(k)).$$

Note that the generic VTK interface for all datasets uses a flat index. The (i, j, k) index can be converted to this flat index as follows: $idx_flat = k \times (npts_x \times npts_y) + j \times nptr_x + i$.

A rectilinear grid consists of cells of the same type. This type is determined by the dimensionality of the dataset (based on the extents) and can either be vertex (0D), line (1D), pixel (2D), or voxel (3D).

Curvilinear grid (structured grid)

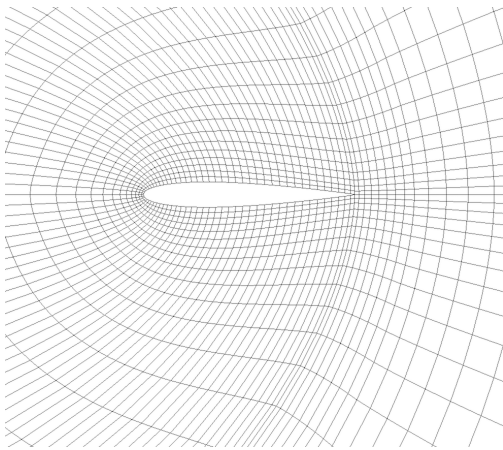


Fig. 1.24: Curvilinear or structured grid.

A curvilinear grid, such as Fig. 1.24, defines its topology implicitly and point coordinates explicitly. To fully define the mesh for a curvilinear grid, VTK uses the following:

- *Extents* - These define the minimum and maximum indices in each direction. For example, a curvilinear grid of extents $(0, 9), (0, 19), (0, 29)$ has $10 \times 20 \times 30$ points regularly defined over a curvilinear mesh.
- *An array of point coordinates* - This array stores the position of each vertex explicitly.

The coordinate of each point is defined as follows: $\text{coordinate} = \text{coordinate_array}(idx_flat)$. The (i, j, k) index can be converted to this flat index as follows: $idx_flat = k \times (npts_x \times npts_y) + j \times npts_x + i$.

A curvilinear grid consists of cells of the same type. This type is determined by the dimensionality of the dataset (based on the extents) and can either be vertex (0D), line (1D), quad (2D), or hexahedron (3D).

AMR dataset

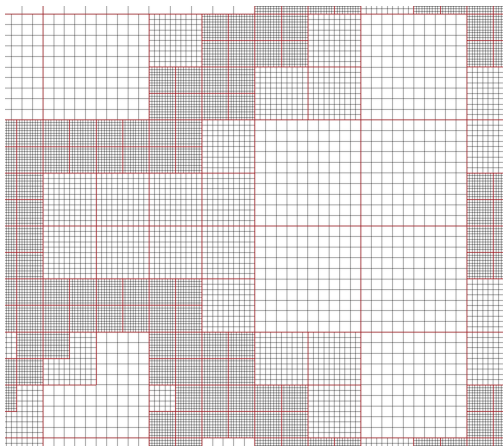


Fig. 1.25: AMR dataset.

VTK natively supports Berger-Oliger type AMR (Adaptive Mesh Refinement) datasets, as shown in Fig. 1.25. An AMR dataset is essentially a collection of uniform rectilinear grids grouped under increasing refinement ratios (decreasing spacing). VTK's AMR dataset does not force any constraint on whether and how these grids should overlap. However,

it provides support for masking (blanking) sub-regions of the rectilinear grids using an array of bytes. This allows VTK to process overlapping grids with minimal artifacts. VTK can automatically generate the masking arrays for Berger-Oliger compliant meshes.

Unstructured grid

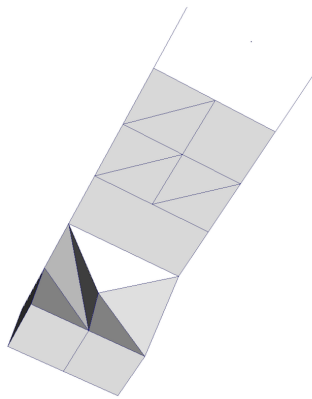


Fig. 1.26: Unstructured grid.

An unstructured grid, such as Fig. 1.26, is the most general primitive dataset type. It stores topology and point coordinates explicitly. Even though VTK uses a memory-efficient data structure to store the topology, an unstructured grid uses significantly more memory to represent its mesh. Therefore, use an unstructured grid only when you cannot represent your dataset as one of the above datasets. VTK supports a large number of cell types, all of which can exist (heterogeneously) within one unstructured grid. The full list of all cell types supported by VTK can be found in the file `vtkCellType.h` in the VTK source code. Here is the list of cell types as of when this document was written:

VTK_EMPTY_CELL	VTK_POLYGON
VTK_VERTEX	VTK_PIXEL
VTK_POLY_VERTEX	VTK_QUAD
VTK_LINE	VTK_TETRA
VTK_POLY_LINE	VTK_VOXEL
VTK_TRIANGLE	VTK_HEXAHEDRON
VTK_TRIANGLE_STRIP	VTK_WEDGE
VTK_PYRAMID	VTK_PARAMETRIC_CURVE
VTK_PENTAGONAL_PRISM	VTK_PARAMETRIC_SURFACE
VTK_HEXAGONAL_PRISM	VTK_PARAMETRIC_TRI_SURFACE
VTK_QUADRATIC_EDGE	VTK_PARAMETRIC_QUAD_SURFACE
VTK_QUADRATIC_TRIANGLE	VTK_PARAMETRIC_TETRA_REGION
VTK_QUADRATIC_QUAD	VTK_PARAMETRIC_HEX_REGION
VTK_QUADRATIC_POLYGON	VTK_HIGHER_ORDER_EDGE
VTK_QUADRATIC_TETRA	VTK_HIGHER_ORDER_TRIANGLE
VTK_QUADRATIC_HEXAHEDRON	VTK_HIGHER_ORDER_QUAD
VTK_QUADRATIC_WEDGE	VTK_HIGHER_ORDER_POLYGON
VTK_QUADRATIC_PYRAMID	VTK_HIGHER_ORDER_TETRAHEDRON
VTK_BIQUADRATIC_QUAD	VTK_HIGHER_ORDER_WEDGE
VTK_TRIQUADRATIC_HEXAHEDRON	VTK_HIGHER_ORDER_PYRAMID
VTK_QUADRATIC_LINEAR_QUAD	VTK_HIGHER_ORDER_HEXAHEDRON
VTK_QUADRATIC_LINEAR_WEDGE	VTK_LAGRANGE_CURVE
VTK_BIQUADRATIC_QUADRATIC_WEDGE	VTK_LAGRANGE_TRIANGLE
VTK_BIQUADRATIC_QUADRATIC_HEXAHEDRON	VTK_LAGRANGE_QUADRILATERAL
VTK_BIQUADRATIC_TRIANGLE	VTK_LAGRANGE_TETRAHEDRON
VTK_CUBIC_LINE	VTK_LAGRANGE_HEXAHEDRON
VTK_CONVEX_POINT_SET	VTK_LAGRANGE_WEDGE
VTK_POLYHEDRON	VTK_LAGRANGE_PYRAMID

Many of these cell types are straightforward. For details, see the VTK documentation.

Polygonal grid (polydata)

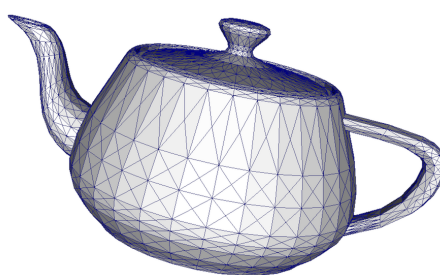


Fig. 1.27: Polygonal grid.

A polydata, such as Fig. 1.27, is a specialized version of an unstructured grid designed for efficient rendering. It consists of 0D cells (vertices and polyvertices), 1D cells (lines and polylines), and 2D cells (polygons and triangle

strips). Certain filters that generate only these cell types will generate a polydata. Examples include the Contour and Slice filters. An unstructured grid, as long as it has only 2D cells supported by polydata, can be converted to a polydata using the `Extract Surface Filter`. A polydata can be converted to an unstructured grid using `Clean to Grid`.

Table

	Author	Affiliation		Alma Mater	Categories	Age	Coolness	
0	Biff	NASA		Ole...	Jazz; Ro...	27	0.6	
1	Bob	Bob's	Supermarket	Ole...	Jazz	54	0.3	
2	Baz	Bob's	Supermarket	TVI	Food	16	0.3	
3	Bippity	Oil	Changes	'R'	TVI	Food	23	0.2
4	Boppity	Oil	Changes	'R'	Home	Food; A...	34	0.25
5	Boo	Oil	Changes	'R'	Princeton	Automobiles	27	0.7

Fig. 1.28: Table

A table, such as Fig. 1.28, is a tabular dataset that consists of rows and columns. All chart views have been designed to work with tables. Therefore, all filters that can be shown within the chart views generate tables. Also, tables can be directly loaded using various file formats such as the comma-separated values format. Tables can be converted to other datasets as long as they are of the right format. Filters that convert tables include `Table to Points` and `Table to Structured Grid`.

Multiblock dataset

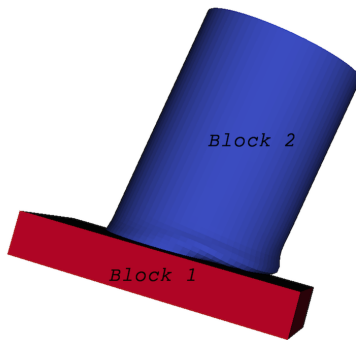


Fig. 1.29: Multiblock dataset.

You can think of a multi-block dataset (Fig. 1.29) as a tree of datasets where the leaf nodes are *simple* datasets. All of the data types described above, except AMR, are *simple* datasets. Multi-block datasets are used to group together datasets that are related. The relation between these datasets is not necessarily defined by **ParaView**. A multi-block dataset can represent an assembly of parts or a collection of meshes of different types from a coupled simulation. Multi-block datasets can be loaded or created within **ParaView** using the `Group` filter. Note that the leaf nodes of a multi-block dataset do not all have to have the same attributes. If you apply a filter that requires an attribute, it will be applied only to blocks that have that attribute.

Multipiece dataset

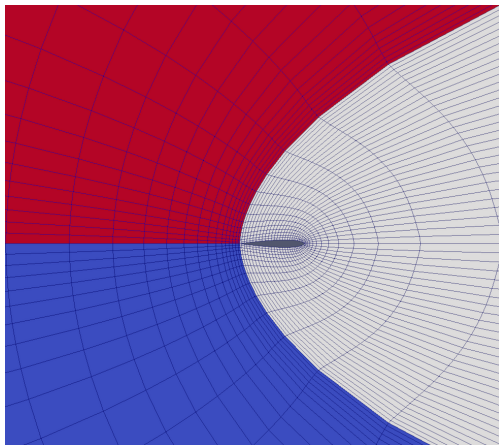


Fig. 1.30: Multipiece dataset.

Multi-piece datasets, such as Fig. 1.30, are similar to multi-block datasets in that they group together simple datasets. There is one key difference. Multi-piece datasets group together datasets that are part of a whole mesh - datasets of the same type and with the same attributes. This data structure is used to collect datasets produced by a parallel simulation without having to append the meshes together. Note that there is no way to create a multi-piece dataset within **ParaView**. It can only be created by using certain readers. Furthermore, multi-piece datasets act, for the most part, as simple datasets. For example, it is not possible to extract individual pieces or to obtain information about them.

1.3.2 Getting data information in paraview

In the visualization pipeline (Section 1.1.2), sources, readers, and filters are all producing data. In a VTK-based pipeline, this data is one of the types discussed. Thus, when you create a source or open a data file in paraview and hit **Apply**, data is being produced. The **Information** panel and the **Statistics Inspector** panel can be used to inspect the characteristics of the data produced by any pipeline module.

The Information panel

The **Information** panel provides summary information about the data produced by the active source. By default, this panel is tucked under a tab below the **Properties** panel. You can toggle its visibility using **View > Information**.

The **Information** panel shows the data information for the active source. Thus, similar to the **Properties** panel, it changes when the active source is changed (e.g., by changing the selection in the **Pipeline Browser**). One way to think of this panel is as a panel showing a summary for the data *currently* produced by the active source. Remember that a newly-created pipeline module does not produce any data until you hit **Apply**. Thus, valid information for a newly-created source will be shown on this panel only after that **Apply**. Similarly, if you change properties on the source and hit **Apply**, this panel will reflect any changes in data characteristics. Additionally, for temporal pipelines, this panel shows information for the current timestep alone (except as noted). Thus, as you step through timesteps in a temporal dataset, the information displayed will potentially change, and the panel will reflect those changes.

Did you know?

Any text on this panel is *copy-able*. For example, if want to copy the number of points value to use it as a property value on the **Properties** panel, simply double-click on the number or click-and-drag to select the number and use the common keyboard shortcut **CTRL + kbd:C** (or **+ C**) to copy that value to the clipboard. Now, you can paste it in an input

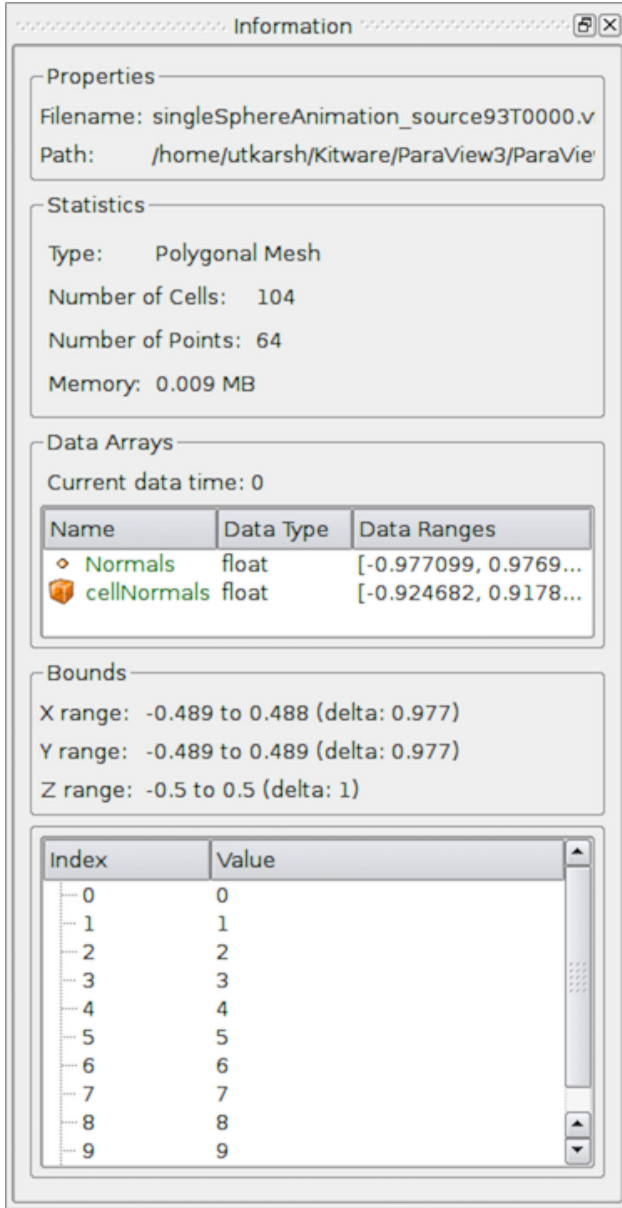


Fig. 1.31: The Information panel in paraview showing data summaries for the active source.

widget in `paraview` or any other application, such as an editor, by using `CTRL + V` (or `⌘ + V`) or the application-specific shortcut for pasting text from the clipboard. The same is true for numbers shown in lists, such as the `Data Ranges`.

The panel itself is comprised of several groups of information. Groups may be hidden based on the type of pipeline module or the type of data being produced.

The file `Properties` group is shown for readers with information about the file that is opened. For a temporal file series, as you step through each time step, the file name is updated to point to the name of the file in the series that corresponds to the current time step.

The `Statistics` group provides a summary of the dataset produced including its type, its number of cells and points (or rows and columns in cases of Tabular datasets), and an estimate of the memory used by the dataset. This number only includes the memory space needed to save the data arrays for the dataset. It does not include the memory space used by the data structures themselves and, hence, must only be treated as an estimate.

The `Data Arrays` group lists all of the available point, cells, and field arrays, as well as their types and ranges for the current time step. The `Current data time` field shows the time value for the current timestep as a reference. As with other places in `paraview`, the icons , , and  are used to indicate cell, point, and field data arrays. Since data arrays can have multiple components, the range for each component of the data array is shown.

`Bounds` shows the spatial bounds of the datasets in 3D Cartesian space. This will be unavailable for non-geometric datasets such as tables.

For reader modules, the `Time` group shows the available time steps and corresponding time values provided by the file.

For structured datasets such as uniform rectilinear grids or curvilinear grids, the `Extents` group is shown that displays the structured extents and dimensions of the datasets.

All of the summary information discussed so far provides a synopsis of the entire dataset produced by the pipeline module, including across all ranks (which will become clearer once we look at using `ParaView` for parallel data processing). In cases of composite datasets, such as multiblock datasets or AMR datasets, recall that these are datasets that are comprised of other datasets. In such cases, these are summaries over all the blocks in the composite dataset. Every so often, you will notice that the `Data Arrays` table lists an array with the suffix `(partial)` (Figure Fig. 1.32). Such arrays are referred to as *partial arrays*. Partial arrays is a term used to refer to arrays that are present on some non-composite blocks or leaf nodes in a composite dataset, but not all. The `(partial)` suffix to indicate partial arrays is also used by `paraview` in other places in the UI.

Name	Data Type	Data Ranges
Normals	float	[-0.974928, 0.974928], [-0...
Result (partial)	double	[0.5, 0.5]

Fig. 1.32: The `Data Arrays` section on `Information` panel showing *partial arrays*. Partial arrays are arrays that present on certain blocks in a composite dataset, but not all.

While summaries over all of the datasets in the composite dataset are useful, you may also want to look at the data information for individual blocks. To do so, you can use the `Data Hierarchy` group, which appears when summarizing composite datasets. The `Data Hierarchy` widget shows the structure or hierarchy of the composite dataset (Figure Fig. 1.33). The `Information` panel switches to showing the summaries for the selected sub-tree. By default, the

root element will be selected. You can now select any block in the hierarchy to view the summary limited to just that sub-tree.

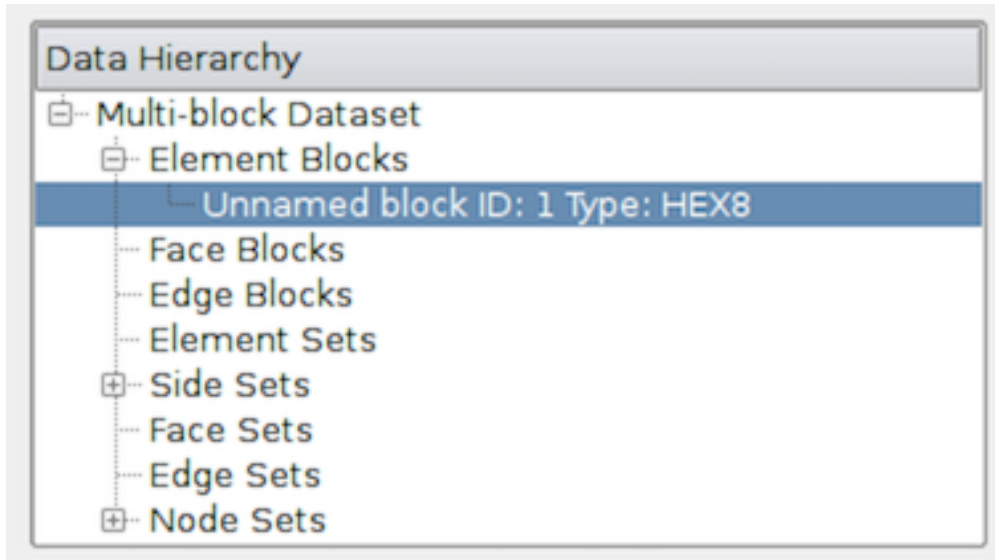


Fig. 1.33: The Data Hierarchy section on the Information panel showing the composite data hierarchy. Selecting a particular block or subtree in this widget will result in the reset of the Information panel showing summaries for that block or subtree alone.

Did you know?

Memory information shown on the Information panel and the Statistics Inspector should only be used as an approximate reference and does not translate to how much memory the data produced by a particular pipeline module takes. This is due to the following factors:

- The size does not include the amount of memory needed to build the data structures to store the data arrays. While, in most cases, this is negligible compared to that of the data arrays, it can be nontrivial, especially when dealing with deeply-nested composite datasets.
- Several filters such as Calculator and Shrink simply pass input data arrays through, so there's no extra space needed for those data arrays that are shared with the input. The memory size numbers shown, however, do not take this into consideration.

If you need an overview of how much physical memory is being used by **ParaView** in its current state, you can use the Memory Inspector (Section 2.7).

The Statistics Inspector panel

The Information panel shows data information for the active source. If you need a quick summary of the data produced by all the pipeline modules, you can use the Statistics Inspector panel. It's accessible from Views > Statistics Inspector.

All of the information on this panel is also presented on the Information panel, except Geometry Size . This corresponds to how much memory is needed for the transformed dataset used for rendering in the active view. For example, to render a 3D dataset as a surface in the 3D view, **ParaView** must extract the surface mesh as a polydata. Geometry Size represents the memory needed for this polydata with the same memory-size-related caveats as with the Information panel.

Statistics Inspector							
Name	Data Type	No. of Cells	No. of Points	Memory (MB)	Geometry Size (MB)	Spatial Bounds	Temporal Bounds
disk_out_ref.ex2	Multi-block Dataset	7472	8499	1.989	0.381	[-5.75, 5.75], [-5.75, 5... [ALL]	
Sphere2	Polygonal Mesh	12480	6242	0.538	0.832	[-0.5, 0.5], [-0.5, 0.5] ... [ALL]	
Wavelet1	Image (Uniform Recti...	8000	9261	0.037	0.002	[-10, 10], [-10, 10], [-10, 10] [ALL]	
AMRGaussianPulse...	Overlapping AMR Da...	605	986	0.02	0.002	[-2, 0.5], [-2, 0.5], [-2, 0.5] [ALL]	
can.ex2	Multi-block Dataset	7152	10088	2.169	1.103	[-7.88, 8.31], [0, 8], [-7.88, 8.31] [0, 0.0043] [ALL]	

Fig. 1.34: The Statistics Inspector panel in paraview showing summaries for all pipeline modules.

1.3.3 Getting data information in pypython

When scripting with **ParaView**, you will often find yourself needing information about the data. While paraview sets up filter properties and color tables automatically using the information from the data, you must do that explicitly when scripting.

In pypython, for any pipeline module (sources, readers, or filters), you can use the following ways to get information about the data produced.

```
>>> from paraview.simple import *
>>> reader = OpenDataFile("../ParaViewData/Data/can.ex2")

# We need to update the pipeline. Otherwise, all of the data
# information we get will be from before the file is actually
# read and, hence, will be empty.
>>> UpdatePipeline()

>>> dataInfo = reader.GetDataInformation()

# To get the number of cells or points in the dataset:
>>> dataInfo.GetNumberOfPoints()
10088
>>> dataInfo.GetNumberOfCells()
7152

# You can always nest the call, e.g.:
>>> reader.GetDataInformation().GetNumberOfPoints()
10088
>>> reader.GetDataInformation().GetNumberOfCells()
7152

# Use source.PointData or source.CellData to get information about
# point data arrays and cell data arrays, respectively.

# Let's print the available point data arrays.
>>> reader.PointData[:]
[Array: ACCL, Array: DISPL, Array: GlobalNodeId, Array: PedigreeNodeId, Array: VEL]

# Similarly, for cell data arrays:
>>> reader.CellData[:]
[Array: EQPS, Array: GlobalElementId, Array: ObjectId, Array: PedigreeElementId]
```

`PointData` (and `CellData`) is a map or dictionary where the keys are the names of the arrays, and the values are objects that provide more information about each of the arrays. In the rest of this section, anything we demonstrate on `PointData` is also applicable to `CellData`.

```
# Let's get the number of available point arrays.  
>>> len(reader.PointData)  
5  
  
# Print the names for all available point arrays.  
>>> reader.PointData.keys()  
['ACCL', 'DISPL', 'GlobalNodeId', 'PedigreeNodeId', 'VEL']  
  
>>> reader.PointData.values()  
[Array: ACCL, Array: DISPL, Array: GlobalNodeId, Array: PedigreeNodeId, Array: VEL]  
  
# To test if a particular array is present:  
>>> reader.PointData.has_key("ACCL")  
True  
  
>>> reader.PointData.has_key("--non-existent-array--")  
False
```

From `PointData` (or `CellData`), you can get access to an object that provides information for each of the arrays. This object gives us methods to get data ranges, component counts, tuple counts, etc.

```
# Let's get information about 'ACCL' array.  
>>> arrayInfo = reader.PointData["ACCL"]  
>>> arrayInfo.GetName()  
'ACCL'  
  
# To get the number of components in each tuple and the number  
# of tuples in the data array:  
>>> arrayInfo.GetNumberOfTuples()  
10088  
>>> arrayInfo.GetNumberOfComponents()  
3  
  
# Alternative way for doing the same:  
>>> reader.PointData["ACCL"].GetNumberOfTuples()  
10088  
>>> reader.PointData["ACCL"].GetNumberOfComponents()  
3  
  
# To get the range for a particular component, e.g. component 0:  
>>> reader.PointData["ACCL"].GetRange(0)  
(-4.965284006175352e-07, 3.212448973499704e-07)  
  
# To get the range for the magnitude in cases of multi-component arrays  
# use -1 as the component number.  
>>> reader.PointData["ACCL"].GetRange(-1)  
(0.0, 1.3329898584157294e-05)  
  
# To determine the data type for this array:  
>>> from paraview import vtk  
>>> reader.PointData["ACCL"].GetDataType() == vtk.VTK_DOUBLE  
True  
# The paraview.vtk module provides access to these constants such as
```

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```
# VTK_DOUBLE, VTK_FLOAT, VTK_INT, etc.

# Likewise, to test the dataset type, itself:
>>> reader.GetDataInformation().GetDataSetType() == \
    vtk.VTK_MULTIBLOCK_DATA_SET
True
```

Here's a sample script to iterate over all point data arrays and print their magnitude ranges:

```
>>> def print_point_data_ranges(source):
...     """Prints array ranges for all point arrays"""
...     for arrayInfo in source.PointData:
...         # get the array's name
...         name = arrayInfo.GetName()
...         # get magnitude range
...         range = arrayInfo.GetRange(-1)
...         print "%s = [%f, %f]" % (name, range[0], range[1])
...
# Let's call this function on our reader.
>>> print_point_data_ranges(reader)
ACCL = [0.000, 0.000]
DISPL = [0.000, 0.000]
GlobalNodeId = [1.000, 10088.000]
PedigreeNodeId = [1.000, 10088.000]
VEL = [0.000, 5000.000]
```

Did you know?

The example scripts in this section all demonstrated how to obtain information about the data such as the number of points and cells, data bounds, and array ranges. However, what they do not show is how to access the raw data itself. To see how to obtain the full data, please see [Section 2.6.11](#).

1.4 Displaying data

The goal of any visualization process is to produce visual representations of the data. The visual representations are shown in modules called views. *Views* provide the canvas on which to display such visual representations, as well as to dictate how these representations are generated from the raw data. The role of the visualization pipeline is often to transform the data so that relevant information can be represented in these views.

Referring back to the visualization pipeline from [Section 1.1.2](#), views are sinks that take in input data but do not produce any data output (i.e., one cannot connect other pipeline modules such as filters to process the results in a view). However, views often provide mechanisms to save the results as images or in other formats including PDF, VRML, and X3D.

Different types of views provide different ways of visualizing data. These can be broadly grouped as follows:

- *Rendering Views* are views that render geometries or volumes in a graphical context. The `Render View` is one such view. Other `Render View`-based views, such as `Slice View` and `Quad View`, extend the basic `render view` to add the ability to add mechanisms to easily inspect slices or generate orthogonal views.
- *Chart Views* cover a wide array of graphs and plots used for visualizing non-geometric data. These include views such as line charts (`Line Chart View`), bar charts (`Bar Chart View`), bag charts (`Bag Chart View`), parallel coordinates (`Parallel Coordinates View`), etc.

- *Comparative Views* are used to quickly generate side-by-side views for parameter study, i.e., to visualize the effects of parameter changes. Comparative variants of **Render View** and several types of the **Chart Views** are available in **ParaView**.

In this chapter, we take a close look at the various views available in **ParaView** and how to use these views for displaying data.

1.4.1 Multiple views

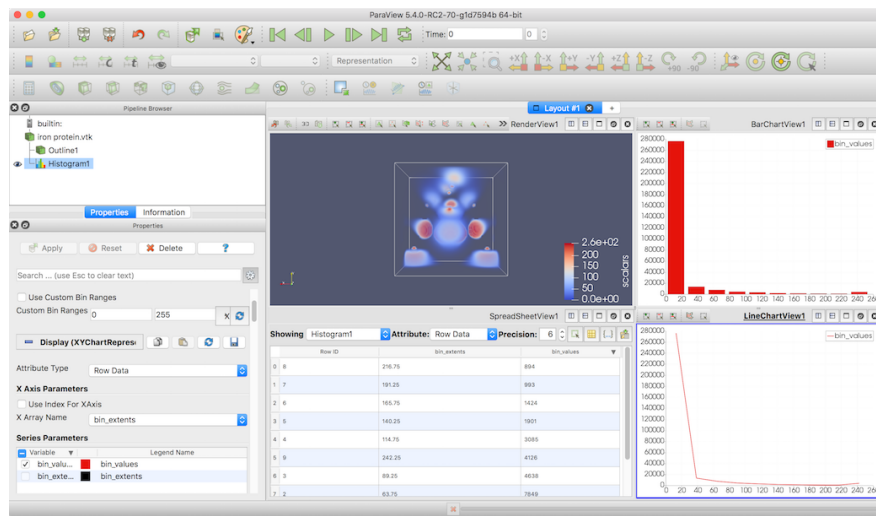


Fig. 1.35: Using multiple views in **paraview** to generate different types of visualizations from a dataset.

With multiple types of views comes the need for creating and viewing multiple views at the same time. In this section, we look at how you can create multiple views and lay them out.

Did you know?

Multiple views were first supported in **ParaView** 3.0. Before that, all data was shown in a single 3D render view, including line plots!

Multiple views in **paraview**

paraview shows all views in the central part of the application window. When **paraview** starts up, the **Render View** is created and shown in the application window by default.

New views can be created by splitting the view frame using the **Split View** controls at the top-right corner of the view frame. Splitting a view splits the view into two equal parts, either vertically or horizontally, based on the button used for the split. On splitting a view, an empty frame with buttons for all known types of views is shown. Simply click on one of those buttons to create a new view of a chosen type.

You can move views by clicking and dragging the title bar for the view (or empty view frame) and dropping it on the title bar on another view (or empty view frame). This will swap the positions of the two views.

Similar to the notion of active source, there is a notion of *active view*. Several panels, toolbars, and menus will update based on the active view. The **Display** properties section on the **Properties** panel, for example, reflects the display properties of the active source in the active view. Similarly, the eyeball icons in the **Pipeline Browser** show the

visibility status of the pipeline module in the active view. Active view is marked in the UI by a blue border around the view frame. Only one view can be active at any time in the application.

Besides being able to create multiple views and laying them out in a pane, paraview also supports placing views in multiple layouts under separate tabs. To create new tabs, use the button in the tab bar. You can close a tab, which will destroy all views laid out in that tab, by clicking on the button. To pop out an entire tab as a separate window, use the button on the tab bar.

The active view is always present in the active tab. Thus, if you change the active tab, the active view will also be changed to be a view in the active tab layout. Conversely, if the active view is changed (by using the Python Shell, for example), the active tab will automatically be updated to be the tab that contains the active view.

Did you know?

You can make the views fullscreen by using *View > Fullscreen*. To return back to the normal mode, use the **Esc** key.

Multiple views in pypython

In pypython, one can create new views using the `CreateView` function or its variants, e.g., `CreateRenderView`.

```
>>> from paraview.simple import *
>>> view = CreateRenderView()
# Alternatively, use CreateView.
>>> view = CreateView("RenderView")
```

When a new view is created, it is automatically made active. You can manually make a view active by using the `SetActiveView` function. Several of the functions available in pypython will use the active view when no view is passed as an argument to the function.

```
# Create a view
>>> view1 = CreateRenderView()
# Create a second view
>>> view2 = CreateRenderView()

# Check if view2 is the active view
>>> view2 == GetActiveView()
True

# Make view1 active
>>> SetActiveView(view1)
>>> view1 == GetActiveView()
True
```

When using Python Shell in paraview, if you create a new view, it will automatically be placed in the active tab by splitting the active view. You can manually control the layout and placement of views from Python too, using the layout API.

In Python, each tab corresponds to a layout.

```
# To get existing tabs/layouts
>>> layouts = GetLayouts()
>>> print(layouts)
{('ViewLayout1', '264'): <paraview.servermanager.ViewLayout object at 0x2e5b7d0>}
```

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```
# To get layout corresponding to a particular view
>>> print(GetLayout(view))
<paraview.servermanager.ViewLayout object at 0x2e5b7d0>

# If view is not specified, active view is used
>>> print(GetLayout())
<paraview.servermanager.ViewLayout object at 0x2e5b7d0>

# To create a new tab
>>> new_layout = servermanager.misc.ViewLayout(registrationGroup="layouts")

# To split the cell containing the view, either horizontally or vertically
>>> view = GetActiveView()
>>> layout = GetLayout(view)
>>> locationId = layout.SplitViewVertical(view=view,
                                             fraction=0.5)
# fraction is optional, if not specified the frame is split evenly.

# To assign a view to a particular cell.
>>> view2 = CreateRenderWindow()
>>> layout.AssignView(locationId, view2)
```

1.4.2 View properties

Just like parameters on pipeline modules, such as readers and filters, views provide parameters that can be used for customizing the visualization such as changing the background color for rendering views and adding title texts for chart views. These parameters are referred to as **View Properties** and are accessible from the **Properties** panel in paraview.

View properties in paraview

Similar to properties on pipeline modules like sources and readers, view properties are accessible from the **Properties** panel. These are grouped under the **View** section. When the active view is changed, the **Properties** panel updates to show the view properties for the active view. Unlike pipeline modules, however, when you change the view properties, they affect the visualization immediately, without use of the **Apply** button.

Did you know?

It may seem odd that **View** and **Display** properties on the **Properties** panel don't need to be **Apply**-ed to take effect, while properties on pipeline modules like sources, readers and filter require you to hit the **Apply** button.

To understand the reasoning behind that, we need to understand why the **Apply** action is needed in the first place. Generally, executing a data processing filter or reader is time consuming on large datasets. If the pipeline module keeps on executing as you are changing the parameter, the user experience will quickly deteriorate, since the pipeline will keep on executing with intermediate (and potentially invalid) property values. To avoid this, we have the **Apply** action. This way, you can set up the pipeline properties to your liking and then trigger the potentially time consuming execution.

Since the visualization process in general focuses on *reducing* data to generate visual representations, the rendering (broadly speaking) is less time- intensive than the actual data processing. Thus, changing properties that affect rendering are not as compute-intensive as transforming the data itself. For example, changing the color on a surface mesh is not

as expensive as generating the mesh in the first place. Hence, the need to **Apply** such properties becomes less relevant. At the same time, when changing display properties such as opacity, you may want to see the result as you change the property to decide on the final value. Hence, it is desirable to see the updates immediately.

Of course, you can always enable **Auto Apply** to have the same immediate update behavior for all properties on the **Properties** panel.

View properties in pypython

In pypython, once you have access to the view, you can directly change view properties on the view object. There are several ways to get access to the view object.

```
# 1. Save reference when a view is created
>>> view = CreateView("RenderView")

# 2. Get reference to the active view.
>>> view = GetActiveView()
```

The properties available on the view will change based on the type of the view. You can use the `help` function to discover available properties.

```
>>> view = CreateRenderView()
>>> help(view)

Help on RenderView in module paraview.servermanager object:

class RenderView(Proxy)
| View proxy for a 3D interactive render
| view.
|
| -----
| Data descriptors defined here:
|
| CenterAxesVisibility
|     Toggle the visibility of the axes showing the center of
|     rotation in the scene.
|
| CenterOfRotation
|     Center of rotation for the interactor.
|
| ...
|
# Once you have a reference to the view, you can then get/set the properties.

# Get the current value
>>> print(view.CenterAxesVisibility)
1

# Change the value
>>> view.CenterAxesVisibility = 0
```

1.4.3 Display properties

Display properties refers to available parameters that control how data from a pipeline module is displayed in a view, e.g., choosing to view the output mesh as a wireframe, coloring the mesh using a data attribute, and selecting which attributes to plot in chart view. A set of display properties is associated with a particular pipeline module and view. Thus, if the data output from a source is shown in two views, there will be two sets of display properties used to control the appearance of the data in each of the two views.

Display properties in paraview

Display properties are accessible from the **Display** section on the **Properties** panel. When the active source or active view changes, this section updates to show the display properties for the active source in the active view, if available. If the active source produces data that cannot be shown (or has never been shown) in the view, then the **Display** properties section may be empty.

Similar to view properties, display property changes are immediately applied, without requiring the use of the **Apply** button.

Display properties in pvython

To access display properties in pvython, you can use `SetDisplayProperties` and `GetDisplayProperty` methods.

```
# Using SetDisplayProperties/GetDisplayProperties to access the display
# properties for the active source in the active view.

>>> print(GetDisplayProperties("Opacity"))
1.0

>>> SetDisplayProperties(Opacity=0.5)
```

Alternatively, you can get access to the display properties object using `GetDisplayProperties` and then changing properties directly on the object.

```
# Get display properties object for the active source in the active view.
>>> disp = GetDisplayProperties()

# You can also save the object returned by Show.
>>> disp = Show()

# Now, you can directly access the properties.
>>> print(disp.Opacity)
0.5

>>> disp.Opacity = 0.75
```

As always, you can use the `help` method to discover available properties on a display object.

```
>>> disp = Show()

>>> help(disp)
>>> help(a)
Help on GeometryRepresentation in module paraview.servermanager object:
```

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```

class GeometryRepresentation(SourceProxy)
| ParaView's default representation for showing any type of
| dataset in the render view.

|
| Method resolution order:
|     GeometryRepresentation
|     SourceProxy
|     Proxy
|     __builtin__.object

-----
| Data descriptors defined here:

|
| ...
|
| CenterStickyAxes
|     Keep the sticky axes centered in the view window.

|
| ColorArrayName
|     Set the array name to color by. Set it to empty string
|     to use solid color.

|
| ColorAttributeType
| ...

```

1.4.4 Render View

Render View is the most commonly used view in **ParaView**. It is used to render geometries and volumes in a 3D scene. This is the view that you typically think of when referring to 3D visualization. The view relies on techniques to map data to graphics primitives such as triangles, polygons, and voxels, and it renders them in a scene.

Most of the scientific datasets discussed in [Section 1.3.1](#) are comprised of meshes. These meshes can be mapped to graphics primitives using several of the established visualization techniques. (E.g., you can compute the outer surface of these meshes and then render that surface as filled polygons, you can just render the edges, or you can render the data as a nebulous blob to get a better understanding of the internal structure in the dataset.) Plugins, like Surface LIC , can provide additional ways of rendering data using advanced techniques that provide more insight into the data.

If the dataset doesn't represent a mesh, e.g., a table ([Section 1.3.1](#)), you cannot directly show that data in this view. However, in such cases, it may be possible to construct a mesh by mapping columns in the table to positions to construct a point cloud, for example.

Understanding the rendering process

Render View uses data processing techniques to map raw data to graphics primitives, which can then be rendered in a 3D scene. These mapping techniques can be classified as follows:

- *Surface rendering* methods provide general rendering by rendering a surface mesh for the dataset. For polygonal datasets ([Section 1.3.1](#)), this is simply the raw data. In cases of other datasets including structured ([Section 1.3.1](#), [Section 1.3.1](#), [Section 1.3.1](#)) and unstructured ([Section 1.3.1](#)) grids, this implies extracting a surface mesh for all external faces in the dataset and then rendering that mesh. The surface mesh itself can then be rendered as a filled surface or as a wireframe simply showing the edges, etc.

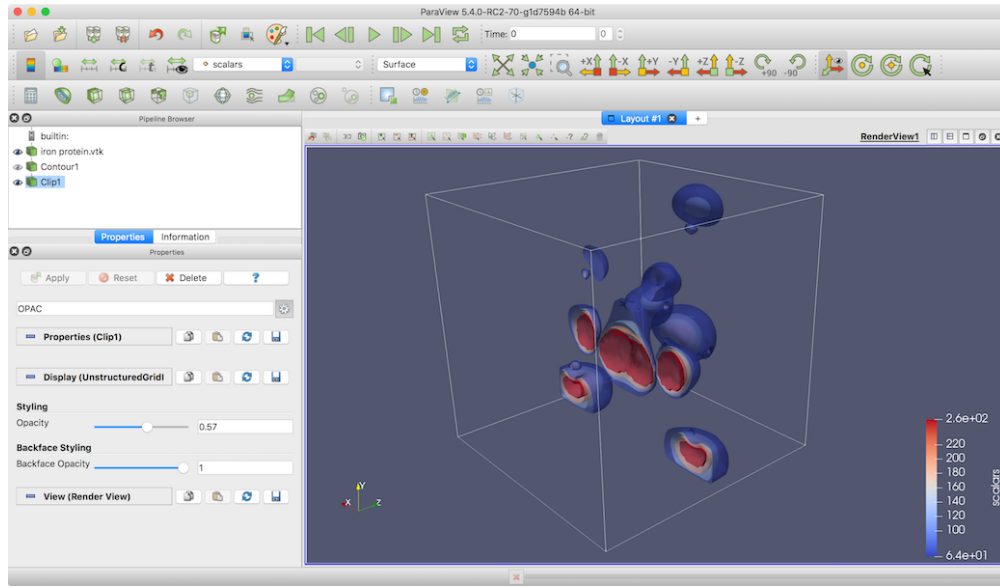


Fig. 1.36: paraview using Render View to generate 3D visualizations from a dataset.

- *Slice* rendering is available for uniform rectilinear grid datasets (Section 1.3.1) where the visualization is generated by simply rendering an orthogonal slice through the dataset. The slice position and alignment can be selected using the display properties.
- *Volume* rendering generates rendering by tracing a ray through the dataset and accumulating intensities based on the color and opacity transfer functions set.

Each of these techniques are referred to as *representations*. When available, you can change the representation type from the display properties on the Properties panel or using the Representation Toolbar .

Render View in paraview

Creating a Render View

Unless you changed the default setting, a new Render View will be created when paraview starts up or connects to a new server. To create a Render View in paraview, split or close a view, and select the Render View button. You can also convert a view to a Render View (or any other type) by right-clicking on the view's title bar and picking from the Convert To sub-menu. It simply closes the chosen view and creates a selected view type in its place.

You can use the Pipeline Browser to control the visibility of datasets produced by pipeline modules in this view. The eyeball icons reflect the visibility state. Clicking on the eyeball icon will toggle the visibility state. If no eyeball icon is shown, it implies that the pipeline module doesn't produce a data type that can be directly shown in the active view, e.g., if the module produced a table, then when Render View is active, there will not be any eyeball icon next to that module.

Interactions

You can interact with the Render View to move the camera in the scene for exploring the visualization and setting up optimal viewing angles. Each of the three mouse buttons, combined with keyboard modifier keys (CTRL or , and), move the camera differently. The interaction mode can be changed from the Camera tab in the **Settings** dialog, which is accessible from *Tools > Settings* (or *ParaView > Preferences* on macOS). There are six interaction modes available in ParaView:

- *Pan* for translating the camera in the view plane.
- *Zoom* for zooming in or out of the center of the view.
- *Roll* for rolling the camera.
- *Rotate* for rotating the camera around the center of rotation.
- *Zoom To Mouse* for zooming in or out of the projected point under the mouse position.
- *Multi Rotate* for allowing azimuth and elevation rotations by dragging from the middle of the view and rolls by dragging from the edges.
- *Rotate Skybox* for rotating the environment skybox. Useful when using Environment Lighting and PBR shader.

The default interactions options are as follows:

Modifier	Left Button	Middle Button	Right Button
	Rotate	Pan	Zoom
	Roll	Rotate	Pan
CTRL or	Rotate Skybox	Rotate	Zoom To Mouse

Usually in ParaView, you are interacting with a 3D scene. However, there are cases when you are working with a 2D dataset such as a slice plane or a 2D image. In such cases, paraview provides a separate set of interaction options suitable for 2D interactions. You can toggle between the default 3D interaction options and 2D interaction options by clicking the 2D or 3D button in the view toolbar. The default interaction options for 2D interactions are as follows:

Modifier	Left Button	Middle Button	Right Button
	Pan	Roll	Zoom
	Zoom	Zoom	Zoom To Mouse
CTRL or	Roll	Pan	Rotate

By default, ParaView will determine whether your data is 2D or 3D when loading the data and will set the interaction mode accordingly. This behavior can be changed in the **Settings** dialog by changing the **Default Interaction Mode** setting under the **Render View** tab. The default setting is “Automatic, based on the first time step”, but the setting can be changed to “Always 2D” or “Always 3D” in case you wish to force the interaction mode.

View properties

Several of the view properties in **Render View** control the annotations shown in the view (Fig. 1.37).

Axes Grid refers to an annotation axis rendered around all datasets in the view (Fig. 1.38). You use the checkbox next to the **Edit Axis Grid** button to show or hide this annotation. To control the annotation formatting, labels, etc., click on the **Edit Axes Grid...** button. The **Axes Grid** is described in Chapter Section 2.10.

The **Center axes** refers to axes rendered in the scene positioned as the center of rotation, i.e., the location is space around which the camera revolves during **Rotate** camera interaction. **Center Axes Visibility** controls the visibility of the center axes.

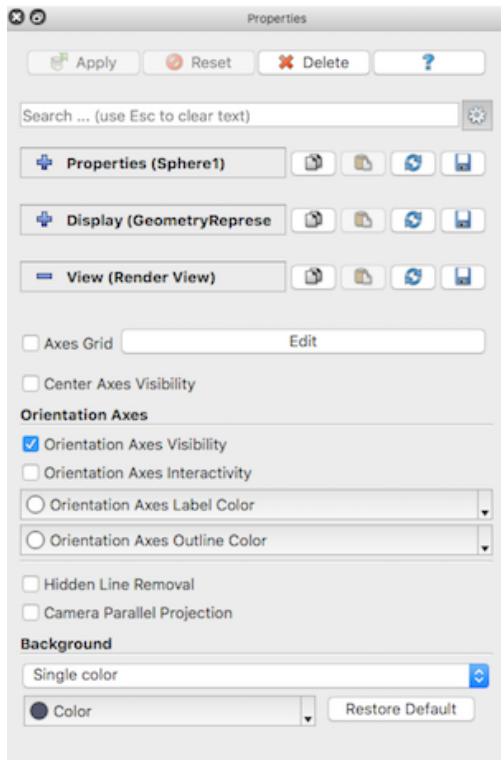


Fig. 1.37: The Properties panel showing view properties for Render View .

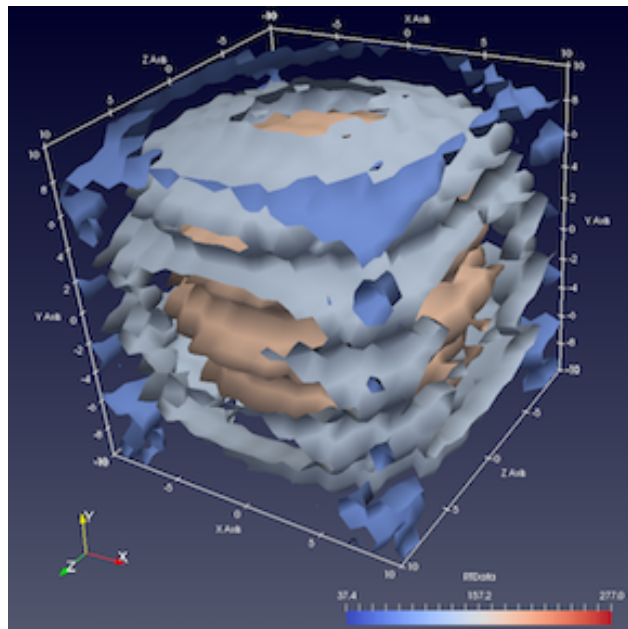


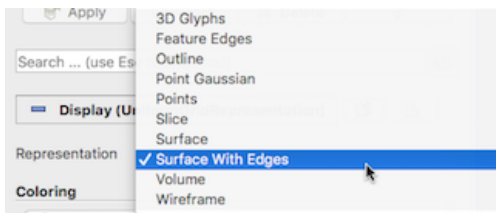
Fig. 1.38: Axes Grid is used to annotate data bounds in Render View .

The *Orientation axes* is the widget shown at the lower-left corner by default, which is used to get a sense for the orientation of the scene. The properties grouped under the **Orientation Axes** group allow you to toggle the visibility and the interactivity of this widget. When interactivity is enabled, you can click and drag this widget to the location of your choosing in the scene.

You can also change the **Background** used for this view. You can either set it as a **Single color** or as a **Gradient** comprised of two colors, or you can select an **Image** (or texture) to use as the background.

There are two advanced properties you may wish to set: hidden line removal and camera parallel projection. The **Hidden Line Removal** option can be enabled to hide lines that would be occluded by a solid object when drawing objects in a **Wireframe** representation. If you want to render your data using parallel projection instead of the default perspective projection you can check the **Camera Parallel Projection** checkbox.

Display properties



One of the first (and probably the most often used) display properties is **Representation**. **Representation** allows you to pick one of the *mapping* modes. The options available depend on the data type, as well as the plugins loaded. While all display properties are accessible from the advanced view for the **Properties** panel, certain properties may be shown/hidden from the default view based on the chosen representation type.

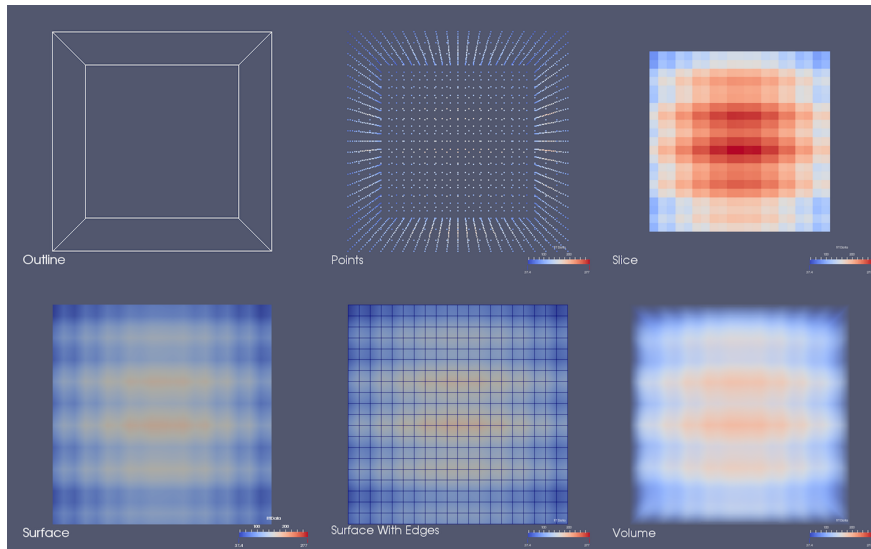


Fig. 1.39: Different renderings generated by rendering data produced by the **Wavelet** source as outline, points, slice, surface, surface with edges, and volume.

The **Outline** representation can be used to render an outline for the dataset. This is arguably the fastest way of rendering the dataset since only the bounding box is rendered. Scalar coloring options, i.e., selecting an array with which to color, has no effect on this representation type. You can still, however, change the **Solid Color** to use as well as the **Opacity**. To change the color, select **Solid Color** in the combo-box under **Coloring**, and then click

Edit to pick the color to use. To change the opacity, simply change the Opacity slider. 0 implies total transparency and, hence, invisiblity, while 1 implies totally opacity.

Did you know?

Rendering translucent data generally adds computational costs to the rendering process. Thus, when rendering large datasets, you may want to leave changing opacities to anything less than 1 to the very end, after having set up the visualization. In doing so, you avoid translucent geometries during exploration, but use them for generating images or screenshots for presentations and publications.

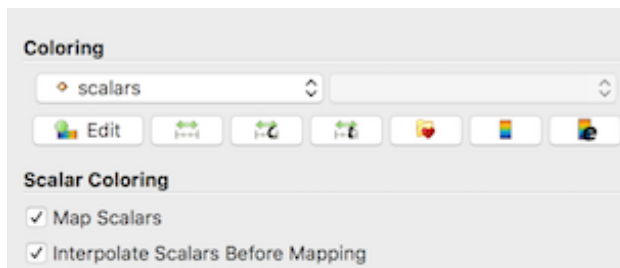
Points , Surface , Surface With Edges , and Wireframe rely on extracting the surface mesh from the dataset and then rendering that either as a collection of points, as solid surface, as solid surface with cell boundaries highlighted, or as a wireframe of cell boundaries only. Feature Edges is a subset of Wireframe consisting of prominent edges on the surface such as edges between cells that form a sharp angle or edges with only one adjacent cell. For these representations, you can either set a single solid color to use, as with Outline , or select a data array to use for scalar coloring (also known as pseudocoloring).

Two other representations are available for most datasets. 3D Glyphs draws a copy of a 3D geometry (e.g., arrow, cone, or sphere, etc.), or glyph, at a subset of points in the dataset. These glyphs can be set to a single color or pseudocolored by a data array. The Point Gaussian representation is similar, but instead of drawing 3D geometry at every point, it draws a 2D image sprite that may have transparency. The image drawn can be one of several predefined image sprites such as Gaussian Blur , Sphere , Black-edged circle , Plain cirlce , Triangle , or Square outline , or a custom sprite can be defined with custom GLSL shader code.

Did you know?

For visualizations that feature 3D glyphs, it is typically much faster to use the 3D Glyph representation rather than the Glyph filter. This is because the glyph representation draws the same geometry at many different locations (a graphics technique called geometry instancing) while the Glyph filter makes many copies of the same source geometry and renders the resulting mesh in its entirety. Generating all the glyphs and rendering them takes potentially a lot of memory and is typically slower to render, so you should use the 3D Glyph representation when possible.

Next, we will cover each of the property groups available under Display properties. Several of these are marked as advanced. Accordingly, you may need to either toggle the panel to show advanced properties using the  button or search for it by name using the search box.



Display properties under Coloring allow you to set how the dataset is colored. To select a single solid color to use to fill the surface or color the wireframe or points, select Solid Color in the combo-box, and then click Edit . That will pop up the standard color chooser dialog from which you can pick a color to use.

If instead you want to pseudocolor using an attribute array available on the dataset, select that array name from the combo-box. For multi-component arrays, you can pick a particular component or Magnitude to use for scalar coloring. ParaView will automatically setup a color transfer function it will use to map the data array to colors. The default range for the transfer function is set up based on the Transfer Function Reset Mode general setting in the Settings

dialog when the transfer function is first created. If another dataset is later colored by a data array with the same name, the range of the transfer function will be updated according to the **Automatic Rescale Range Mode** property in the **Color Map Editor**. To reset the transfer function range to the range of the data array in the selected dataset, you can use the **Rescale** button. Remember that, despite the fact that you can set the scalar array with which to color when rendering as **Outline**, the outline itself continues to use the specified solid color.

Scalar Coloring properties are only relevant when you have selected a data array with which to pseudocolor. The **Map Scalars** checkbox affects whether a color transfer function should be used (Fig. 1.40). If unchecked, and the data array can directly be interpreted as colors, then those colors are used directly. If not, the color transfer function will be used. A data array can be interpreted as colors if, and only if, it is an unsigned char, float, or double array with two, three, or four components. If the data array is unsigned char, the color values are defined between 0 and 255 while if the data array is float or double, the color values are expected to be between 0 and 1. **Interpolate Scalars Before Mapping** controls how color interpolation happens across rendered polygons. If on, scalars will be interpolated within polygons, and color mapping will occur on a per-pixel basis. If off, color mapping occurs at polygon points, and colors are interpolated, which is generally less accurate. Refer to the Kitware blog [PatMarion] for a detailed explanation of this option.

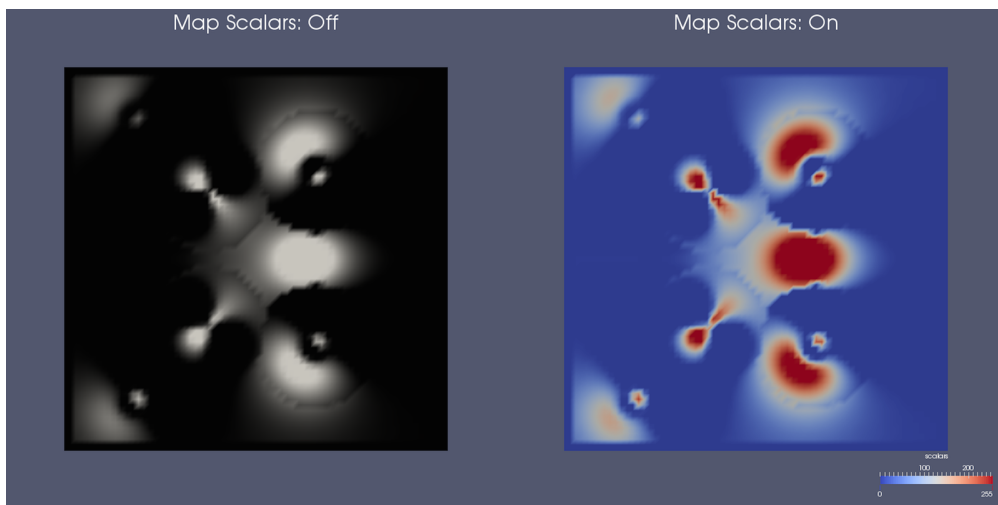
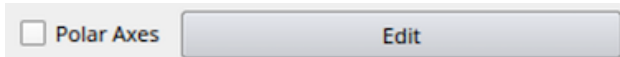


Fig. 1.40: The **Map Scalars** property can be used to avoid using a transfer function and directly interpreting the array values as colors, if possible.



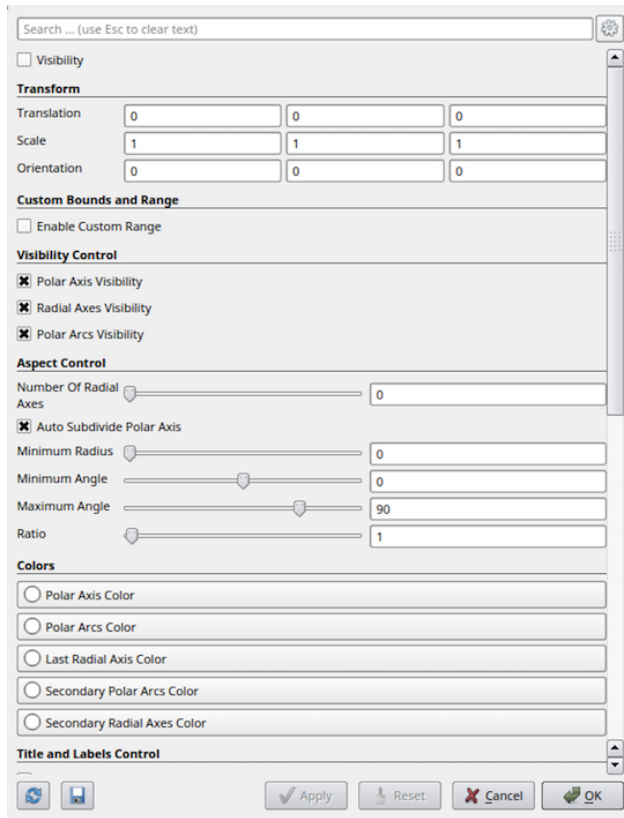
The **Polar Axes** checkbox toggles polar axes shown around the data. Many parameters can be accessed via an **Edit** button alongside it. The parameters include angles, tick range, labels, logarithmic mode, ellipse ratio and more.

Styling properties include **Opacity** (useful when rendering translucent geometries), **Point Size** (used to control size of points rendered with using **Points** representation), and **Line Width** (used to control the thickness of lines when rendering as **Wireframe** or that of the edges when rendering as **Surface With Edges**).

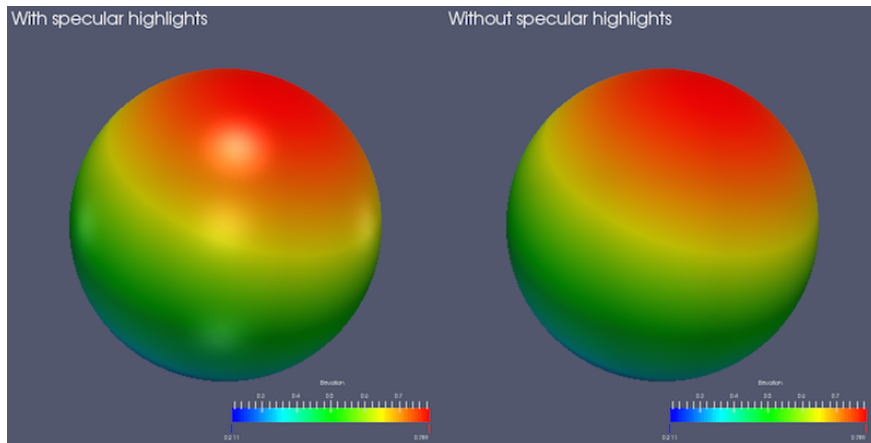
Lighting properties affect the shading for rendered surfaces. **Interpolation** allows you to pick between **Flat** and **Gouraud** shading. **Specular**, together with **Specular Color** and **Specular Power**, affects the shininess of the surface. Set this to a non-zero value to render shiny, metallic surfaces.

Common Errors

Specular highlights can lead to misinterpretation of scalar values when using scalar coloring, since the color shown on the shiny part of the surface will not correspond to any color on the color transfer function. Hence, it is generally



advisable to use specular highlights on surfaces colored with a single solid color and not on those using scalar coloring (or pseudocoloring).



Edge Styling allows you to set the Edge Color with which to color the edges when using Surface With Edges representation.

Backface Styling provides advanced controls to fine-tune the rendering by controlling front and back faces. A front face is any mesh face facing the camera, while a back face is the one facing away from the camera. By choosing to Cull Frontface or Cull Backface, or by selecting a specific representation type to use for the backface, you can customize your visualizations.

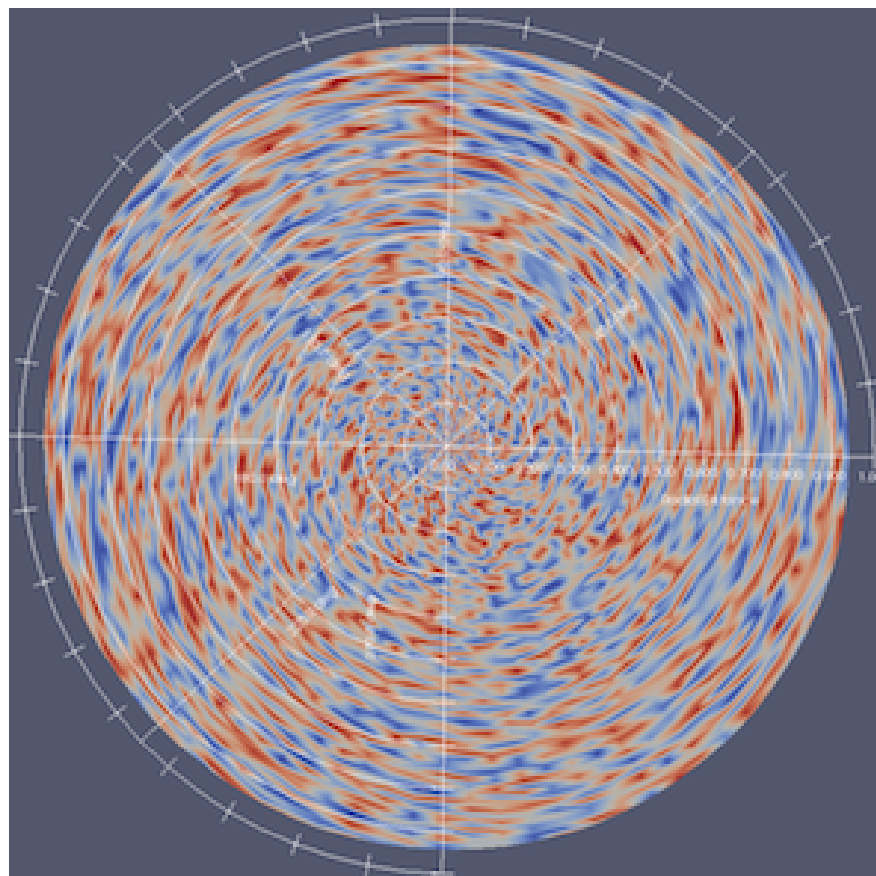


Fig. 1.41: A Polar Axes usage example.

Styling

Opacity 1

Point Size

Line Width

Lighting

Interpolation Gouraud

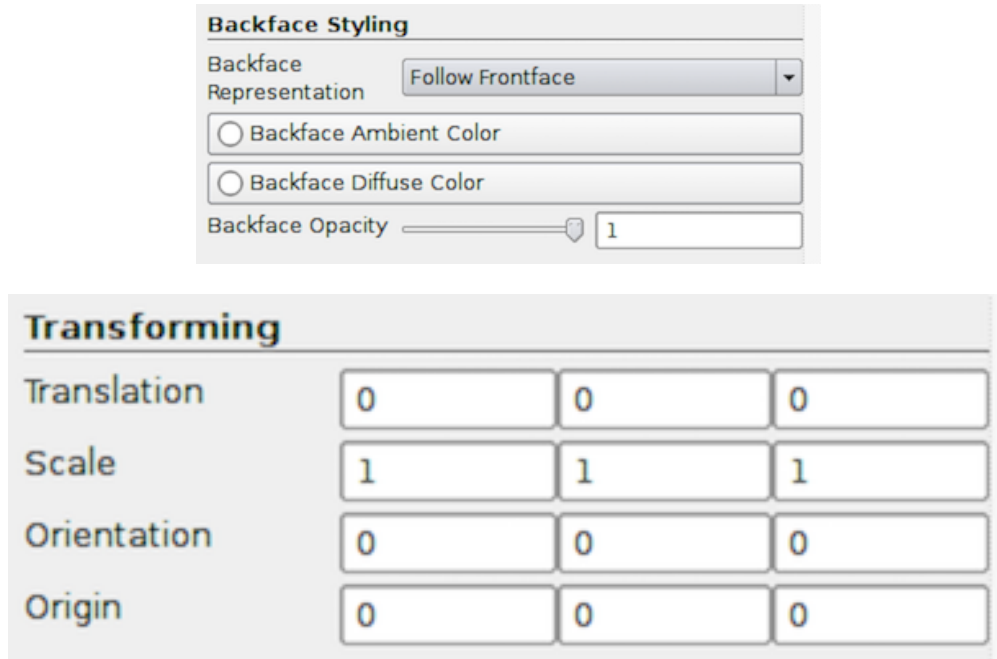
Specular 0

Specular Color

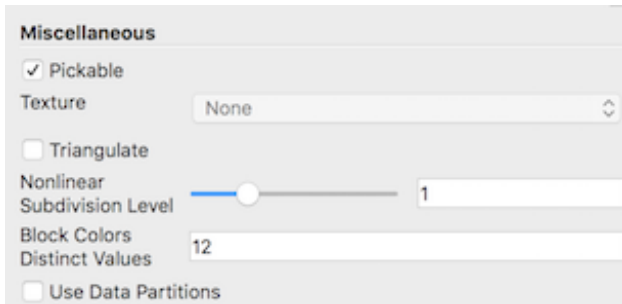
Specular Power 100

Edge Styling

Edge Color



Transforming properties can be used to transform the rendered data in the scene without affecting the raw data itself. Thus, if you apply filters on the data source, it will indeed be working with the untransformed data. To transform the data itself, you should use the **Transform** filter.



Several properties are available under the **Miscellaneous** group. Uncheck the **Pickable** option if you want the dataset to be ignored when making selections. If the dataset has a texture coordinates array, you can apply a texture to the dataset surface using the **Texture** combo-box. Choose **Load** to load a texture or apply a previously loaded texture listed in the combo-box. If your dataset doesn't have texture coordinates, you can create them by applying one of **Texture Map to Cylinder**, **Texture Map to Sphere**, or **Texture Map To Plane** filters, or using the filters **Calculator** or **Programmable Filter**.

The **Triangulate** option is useful for rendering objects with non-convex polygons. It comes with some additional processing cost for converting polygons to triangles, so it should be used only when necessary.

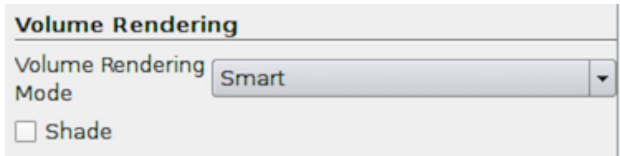
The **Nonlinear Subdivision Level** is used when rendering datasets with higher-order elements. Use this to set the subdivision level for triangulating higher order elements. The higher the value, the smoother the edges. This comes at the cost of more triangles and, hence, potentially, increased rendering time.

The **Block Colors Distinct Values** sets the number of unique colors to use when coloring multi-block datasets by block ID. Finally, **Use Data Partitions** controls whether data is redistributed when it is rendered translucently. When off (default value), data is repartitioned by the compositing algorithm prior to rendering. This is typically an expensive operation that slows down rendering. When this option is on, the existing data partitions are used, and the cost of data redistribution is avoided. However, if the partitions are not sortable in back-to-front order, rendering artifacts

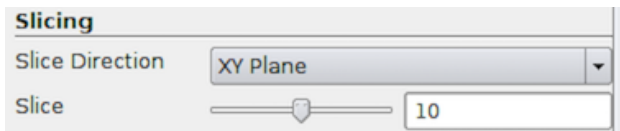


Fig. 1.42: A dataset made of quadratic tetra hedra displayed with 1, 2, and 3 levels of nonlinear subdivision.

may occur.



Volume Rendering options are available if the data can be volume rendered. You can pick a specific type of Volume Rendering Mode , although the default (Smart) should work in most cases, since it attempts to pick a volume rendering mode suitable for your data and graphics setup. To enable gradient-based shading, check Shade , if available.



Slicing properties are available when the Slice representation type is present. These allow you to pick the orthogonal slice plane orientation and slice offset using Slice Direction and the Slice slider.

Render View in pvpthon

Creating a Render View

You use CreateRenderView or CreateView functions to create a new instance of a render view.

```
>>> from paraview.simple import *
>>> view = CreateRenderView()
# Alternatively, use CreateView.
>>> view = CreateView("RenderView")
```

You use Show and Hide to show or hide data produced by a pipeline module in the view.

```
>>> source = Sphere()
>>> view = CreateRenderView()

# Show active source in active view.
>>> Show()

# Or specify source and view explicitly.
>>> Show(source, view)

# Hide source in active view.
>>> Hide(source)
```

Interactions

Since pvvpython is designed for scripting and batch processing, it has limited support for direct interaction with the view. To interact with a scene, invoke the `Interact` function in Python.

Interact()

More often, you will programmatically change the camera as follows:

```
# Get camera from the active view, if possible.  
>>> camera = GetActiveCamera()  
  
# or, get the camera from a specific render view.  
>>> camera = view.GetActiveCamera()  
  
# Now, you can use methods on camera to move it around the scene.  
  
# Divide the camera's distance from the focal point by the given dolly value.  
# Use a value greater than one to dolly-in toward the focal point, and use a  
# value less than one to dolly-out away from the focal point.  
>>> camera.Dolly(10)  
  
# Set the roll angle of the camera about the direction of projection.  
>>> camera.Roll(30)  
  
# Rotate the camera about the view up vector centered at the focal point. Note  
# that the view up vector is whatever was set via SetViewUp, and is not  
# necessarily perpendicular to the direction of projection. The result is a  
# horizontal rotation of the camera.  
>>> camera.Azimuth(30)  
  
# Rotate the focal point about the view up vector, using the camera's position  
# as the center of rotation. Note that the view up vector is whatever was set  
# via SetViewUp, and is not necessarily perpendicular to the direction of  
# projection. The result is a horizontal rotation of the scene.  
>>> camera.Yaw(10)  
  
# Rotate the camera about the cross product of the negative of the direction  
# of projection and the view up vector, using the focal point as the center  
# of rotation. The result is a vertical rotation of the scene.  
>>> camera.Elevation(10)  
  
# Rotate the focal point about the cross product of the view up vector and the  
# direction of projection, using the camera's position as the center of  
# rotation. The result is a vertical rotation of the camera.  
>>> camera.Pitch(10)
```

Alternatively, you can explicitly set the camera position, focal point, view up, etc., to explicitly place the camera in the scene.

```
>>> camera.SetFocalPoint(0, 0, 0)  
>>> camera.SetPosition(0, 0, -10)  
>>> camera.SetViewUp(0, 1, 0)
```

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```
>>> camera.SetViewAngle(30)
>>> camera.SetParallelProjection(False)

# If ParallelProjection is set to True, then you'll need
# to specify parallel scalar as well i.e. the height of the viewport in
# world-coordinate distances. The default is 1. Note that the 'scale'
# parameter works as an 'inverse scale' where larger numbers produce smaller
# images. This method has no effect in perspective projection mode.
>>> camera.SetParallelScale(1)
```

View properties

In pypython, view properties are directly accessible on the view object returned by `CreateRenderWindow` or `GetActiveView`.

Once you get access to the view properties objects, you can then set properties on it similar to properties on pipeline modules such as sources, filters, and readers.

```
>>> view = GetActiveView()

# Set center axis visibility
>>> view.CenterAxesVisibility = 0

# Or you can use this variant to set the property on the active view.
>>> SetViewProperties(CenterAxesVisibility=0)

# Another way of doing the same
>>> SetViewProperties(view, CenterAxesVisibility=0)

# Similarly, you can change orientation axes related properties
>>> view.OrientationAxesVisibility = 0
>>> view.OrientationAxesLabelColor = (1, 1, 1)
```

Display properties

Similar to view properties, display properties are accessible from the display properties object or using the `SetDisplayProperties` function.

```
>>> displayProperties = GetDisplayProperties(source, view)
# Both source and view are optional. If not specified, the active source
# and active view will be used.

# Now one can change properties on this object
>>> displayProperties.Representation = "Outline"

# Or use the SetDisplayProperties API.
>>> SetDisplayProperties(source, view, Representation=Outline)

# Here too, source and view are optional and when not specified,
# active source and active view will be used.
```

You can always use the `help` function to get information about available properties on a display properties object.

1.4.5 Line Chart View

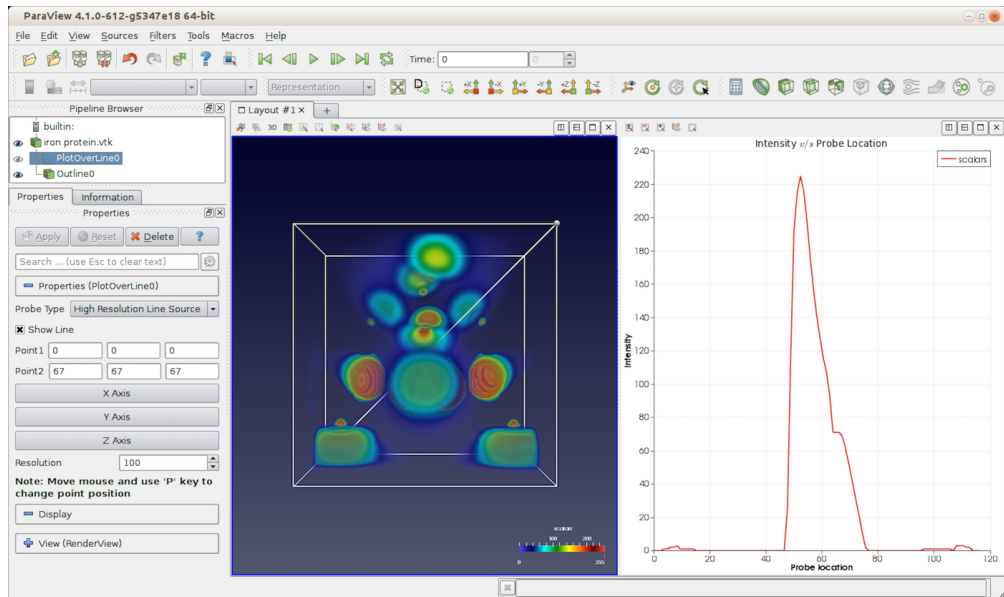


Fig. 1.43: `paraview` using `Line Chart View` to plot data values probed along a line through the dataset using `Plot Over Line` filter.

`Line Chart View` can be used to plot data as a line plot representing changes in dependent variables against an independent variable. Using display properties, you can also show scatter plots in this view. This view and other charting views in **ParaView** follow a similar design, where you pick attribute arrays to plot using display properties, and they are plotted in the view. How those values get plotted depends on the type of the view: `Line Chart View` draws a line connecting sample points, `Bar Chart View` renders bars at each sample point, etc.

One of the most common ways of showing a line plot is to apply the `Plot Over Line` filter to any dataset. This will probe the dataset along the probe line specified. You then plot the sampled values in the `Line Chart View`. Alternatively, if you have a tabular dataset (i.e. `vtkTable`), then you can directly show the data in this view.

Did you know?

You can plot any arbitrary dataset, even those not producing `vtkTable` outputs, by using the `Plot Data` filter. Remember, however, that for extremely large datasets, while `Render View` may use parallel rendering strategies to improve performance and reduce memory requirements, chart views rarely, if ever, support such parallel strategies.

Understanding plotting

Line Chart View plots data arrays. For any dataset being shown in the view, you first select which data array is to be treated as the independent variable and plotted along the x-axis. Then, you select which arrays to plot along the Y-axis. You can select multiple of these and setup properties for each of the series so they are rendered with different colors and line styles. Since data arrays in VTK datasets are associated with cells or points, and the two are not directly comparable to one another, you can only pick arrays associated with one type of attribute at any time.

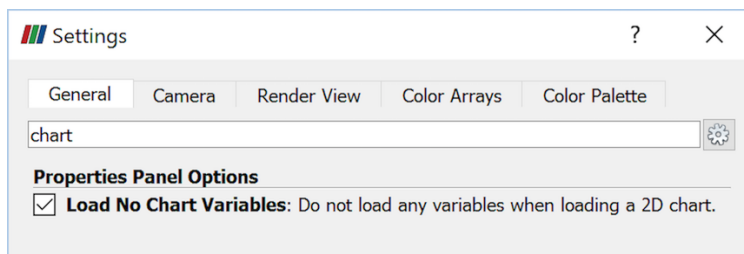
Line Chart View in paraview

Creating a Line Chart View

Similar to creating **Render View**, you can split the viewport or convert an existing view to **Line Chart View**. **Line Chart View** will also be automatically created if you apply a filter that needs this view, e.g., the **Plot Over Line** filter.

Did you know?

If you generate lengthy data for the **Line Chart View**, the default variables that are selected may be slow to adjust. You can change **paraview**'s default behavior to initially load no variables at all by selecting the **Load No Chart Variables** checkbox under **Settings/General/Properties Panel Options**.



Interactions

Interactions with the chart view result in changing the plotted axes ranges. You can left-click and drag to pan, i.e., change the origin. To change the range on either of the axes, you can right-click and drag vertically and/or horizontally to change the scale on the vertical axes and/or horizontal axes, respectively.

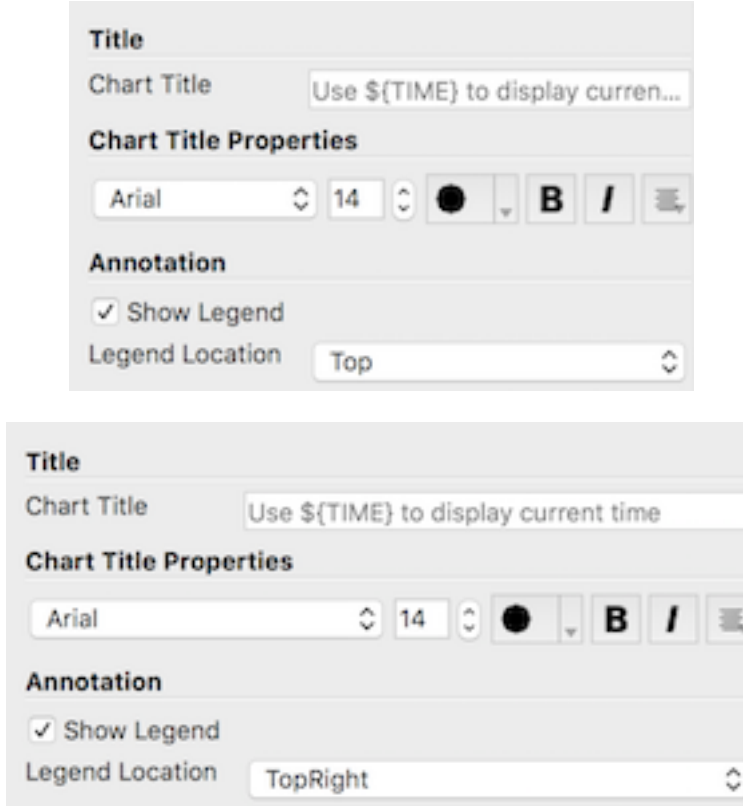
You can also explicitly specify the axes range using view properties.

View properties

The view properties for **Line Chart View** are grouped as properties that affect the view and those that affect each of the potential four axes.

To set a title, use **Chart Title**. Title text properties such as font, size, style, and alignment with respect to the chart can be set under **Chart Title Properties**. To toggle the visibility of the legend, use **Show Legend**. While you cannot interactively place the legend in this view, you can use **Legend Location** to place it at one of the corners.

There are four axes possible in this view: left, bottom, top, and right. The top and right axes are shown only when some series is set to use those. (We will cover this in the *Display properties* subsection.) For each of the axes, you can



set a title (e.g., `Left Axis Title`, `Bottom Axis Title`, etc.) and adjust the title font properties. You can turn on a grid with a customizable color by checking the `Show Left Axis Grid`, for example.

Next, you can customize the axes ranges. You can always simply interact with the mouse to set the axes ranges. To precisely set the range, check the `Axis Use Custom Range` for the appropriate axis, e.g., `Bottom Axis Use Custom Range` for fixing the bottom axis range, and then specify the data values to use for the min and the max.

The labels on the axes are, by default, automatically determined to avoid visual clutter. By default, the axis labels are arranged on a linear scale, but by enabling the `Axis Log Scale` option you can use log scaling instead. In addition, you can override the default labelling strategy for any of the axes separately and, instead, specify the locations to label explicitly. This can be done by checking `Axis Use Custom Labels` for a particular axis, e.g., `Bottom Axis Use Custom Labels`. When checked, a list widget will be shown where you can manually add values at which labels will be placed.

For generating log plots, simply check the corresponding `Axis Use Log Scale`, e.g., `Left Axis Use Log Scale` to use log scale for Y-axis (Fig. 1.44). Note that log scale should only be used for an axis with a non-zero positive range, since the log of a number less than or equal to 0 is undefined.

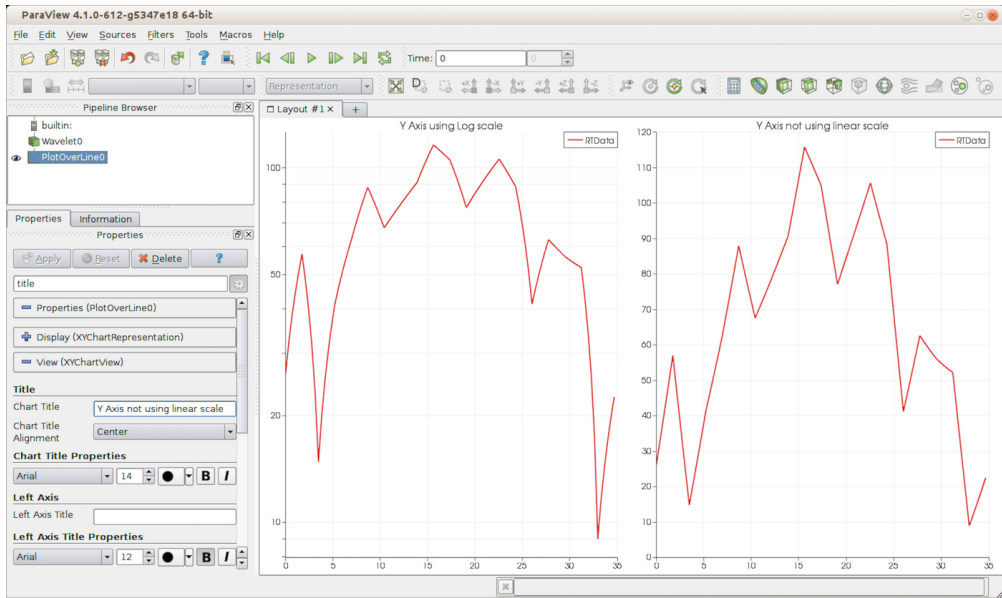
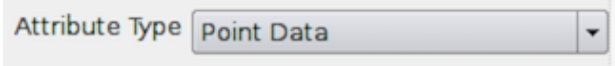
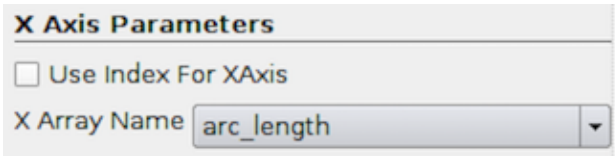


Fig. 1.44: Differences between line charts when using log scale for the Y-axis.

Display properties



Display properties allow you to setup which series or data arrays are plotted in this view. You start by picking the **Attribute Type**. Select the attribute type that has the arrays of interest. For example, if you are plotting arrays associated with points, then you should pick **Point Data**.) Arrays with different associations cannot be plotted together. You may need to apply filters such as **Cell Data to Point Data** or **Point Data to Cell Data** to convert arrays between different associations for that.

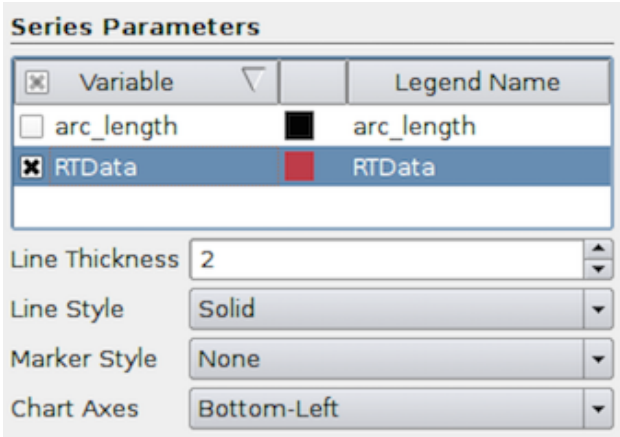


Properties under **X Axis Parameters** allow you to select the independent variable plotted on the X axis by choosing the **X Array Name**. If none of the arrays are appropriate, you can choose to use the element index in the array as the X axis by checking **Use Index for XAxis**.

Series Parameters control series or data arrays plotted on the Y-axis. All available data arrays are lists in the table widget that allows you to check/uncheck a series to plot in the first column. The second column in the table shows the associated color used to plot that series. You can double-click the color swatch to change the color to use. By default, **ParaView** will try to pick a palette of discrete colors. The third column shows the label to use for that series in the legend. By default, it is set to be the same as the array name. You can double-click to change the name to your choice, e.g., to add units.

Other series parameters include **Line Thickness**, **Line Style**, and **Marker Style**. To change any of these, highlight a row in the **Series Parameters** widget, and then change the associated parameter to affect the highlighted series. You can change properties for multiple series and can select multiple of them by using the **CTRL** (or **Shift**) keys.

Using **Chart Axes**, you can change which axes on which a series is shown. The default is **Bottom-Left**, but you



can change it to be `Bottom-Right`, `Top-Left`, or `Top-Right` to accommodate series with widely different ranges in the same plot.

Line Chart View in pvython

The principles involved in accessing `Line Chart View` from pvython are similar to those with `Render View`. You work with the view properties and display properties objects to change views and display properties, respectively. The thing that changes is the set of available properties.

The following script demonstrates the typical usage:

```
>>> from paraview.simple import *

# Create a data source to probe into.
>>> Wavelet()
<paraview.servermanager.Wavelet object at 0x1156fd810>

# We update the source so that when we create PlotOverLine filter
# it has input data available to determine good defaults. Otherwise,
# we will have to manually set up the defaults.
>>> UpdatePipeline()

# Now, create the PlotOverLine filter. It will be initialized using
# defaults based on the input data.
>>> PlotOverLine()
<paraview.servermanager.PlotOverLine object at 0x1156fd490>

# Show the result.
>>> Show()
<paraview.servermanager.XYChartRepresentation object at 0x1160a6a10>

# This will automatically create a new Line Chart View if the
# the active view is no a Line Chart View since PlotOverLine
# filter indicates it as the preferred view. You can also explicitly
# create it by using CreateView() function.

# Display the result.
>>> Render()
```

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```
# Access display properties object.
>>> dp = GetDisplayProperties()
>>> print(dp.SeriesVisibility)
['arc_length', '0', 'RTData', '1']

# This is list with key-value pairs where the first item is the name
# of the series, then its visibility and so on.
# To toggle visibility, change this list e.g.
>>> dp.SeriesVisibility = ['arc_length', '1', 'RTData', '1']

# Same is true for other series parameters including series color,
# line thickness etc.

# For series color, the value consists of 3 values: red, green, and blue
# color components.
>>> print(dp.SeriesColor)
['arc_length', '0', '0', '0', 'RTData', '0.89', '0.1', '0.11']

# For series labels, value is the label to use.
>>> print(dp.SeriesLabel)
['arc_length', 'arc_length', 'RTData', 'RTData']

# e.g. to change RTData's legend label, we can do something as follows:
>>> dp.SeriesLabel[3] = 'RTData -- new label'

# Access view properties object.
>>> view = GetActiveView()
# or
>>> view = GetViewProperties()

# To change titles
>>> view.ChartTitle = "My Title"
>>> view.BottomAxisTitle = "X Axis"
>>> view.LeftAxisTitle = "Y Axis"
```

1.4.6 Bar Chart View

Bar Chart View is very similar to Line Chart View when it comes to creating the view, view properties, and display properties. One difference is that, instead of rendering lines for each series, this view renders bars. In addition, under the display properties, the Series Parameters like Line Style and Line Thickness are not available, since they are not applicable to bars.

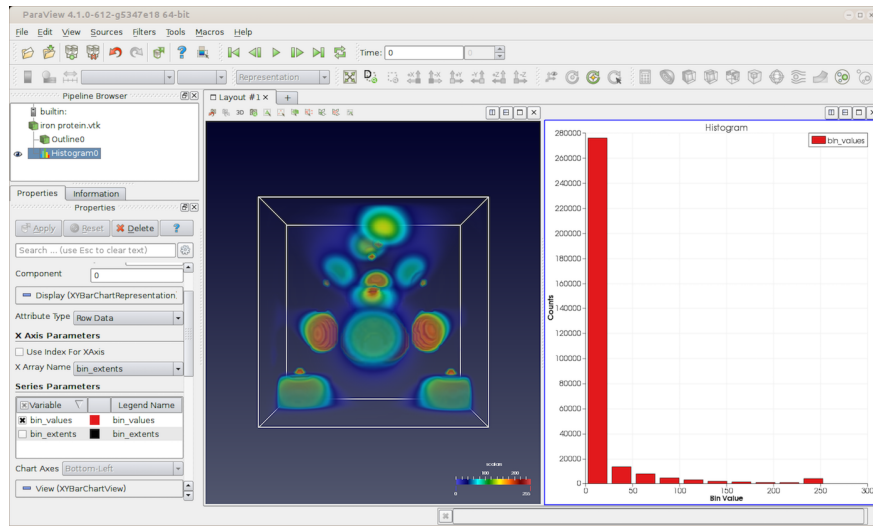


Fig. 1.45: paraview using Bar Chart View to plot the histogram for a dataset using the Histogram filter.

1.4.7 Box Chart View

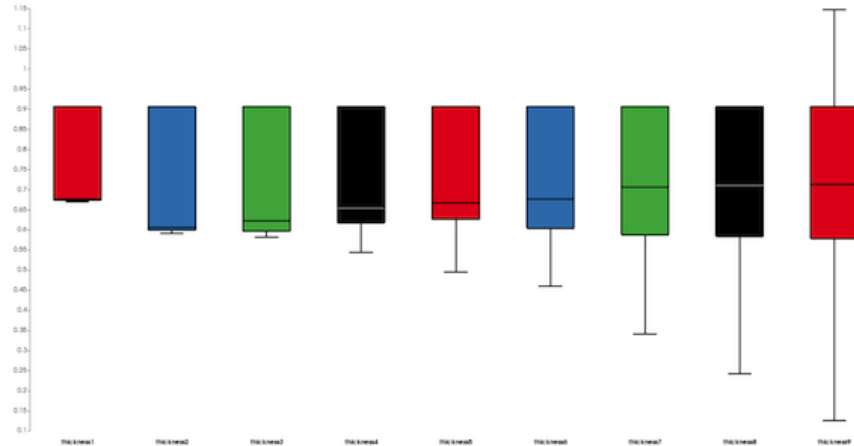


Fig. 1.46: paraview using Box Chart View to plot the box chart for a dataset using the Compute Quartiles filter.

Box plot is a standard method for graphically depicting groups of statistical data based on their quartiles. A box plot is represented by a box with the following properties: the bottom of the rectangle corresponds to the first quartile, a horizontal line inside the rectangle indicates the median and the top of the rectangle corresponds to the third quartile. The maximum and minimum values are depicted using vertical lines that extend from the top and the bottom of the rectangle.

In ParaView, the Box Chart View can be used to display such box plots through the Compute Quartiles filter which computes the statistical data needed by the view to draw the box plots.

1.4.8 Plot Matrix View

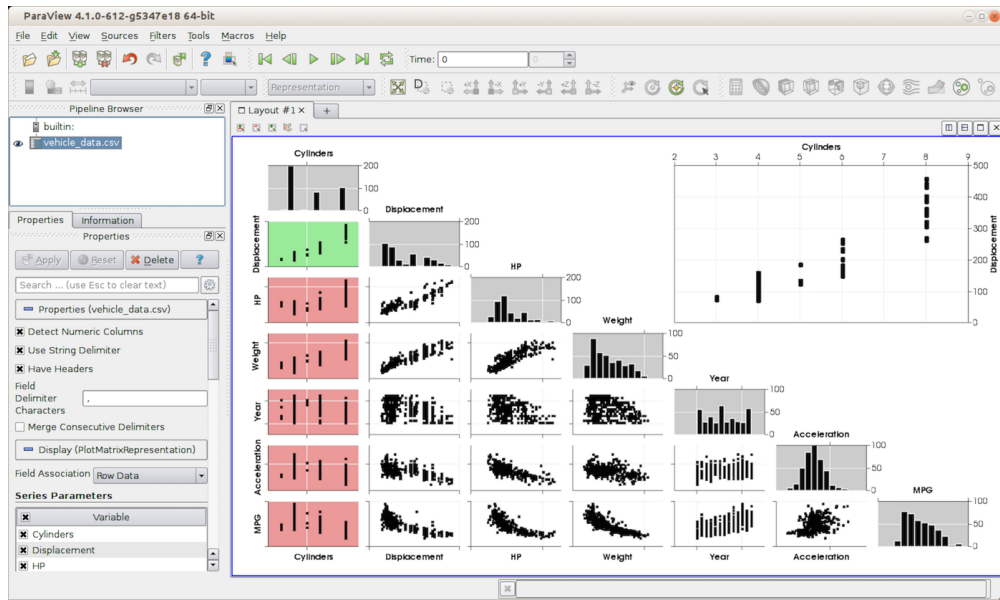


Fig. 1.47: paraview using Plot Matrix View to generate a scatter plot matrix to understand correlations between pairs of variables.

Plot Matrix View is a chart view that renders a scatter plot matrix. It allows you to spot patterns in the small scatter plots, change focus to those plots of interest, and perform basic selection. The principle is that, for all selected arrays or series to be plotted, the view generates a scatter plot for each pair. You can activate a particular scatter plot, in which case the active plot is re-drawn at a bigger scale to make it easier to inspect. Additionally, the view shows a histogram for each plotted variable or series.

The view properties allow you to set colors to use for active plot, histograms, etc., while the display properties allow you to pick which series are plotted.

Interactions

You can click on any of the plots (except the histograms) in the matrix to make it active. Once activated, the active plot will show that plot. You can then interact with the active plot exactly like Line Chart View or Bar Chart View for panning and zoom.

View properties

View properties on this view allow you to pick styling parameters for the rendering ranging from title (Chart Title) to axis colors (Active Plot Axis Color, Active Plot Grid Color). You can also control the visibility of the histogram plots, the active plot, the axes labels, the grids, and so on.

Display properties

Similar to Line Chart View, you select the Attribute Type and then the arrays to plot. Since, in a scatter plot matrix, the order in which the selected series are rendered can make it easier to understand correlations, you can change the order by clicking and dragging the rows in the Series Parameters table.

1.4.9 Parallel Coordinates View

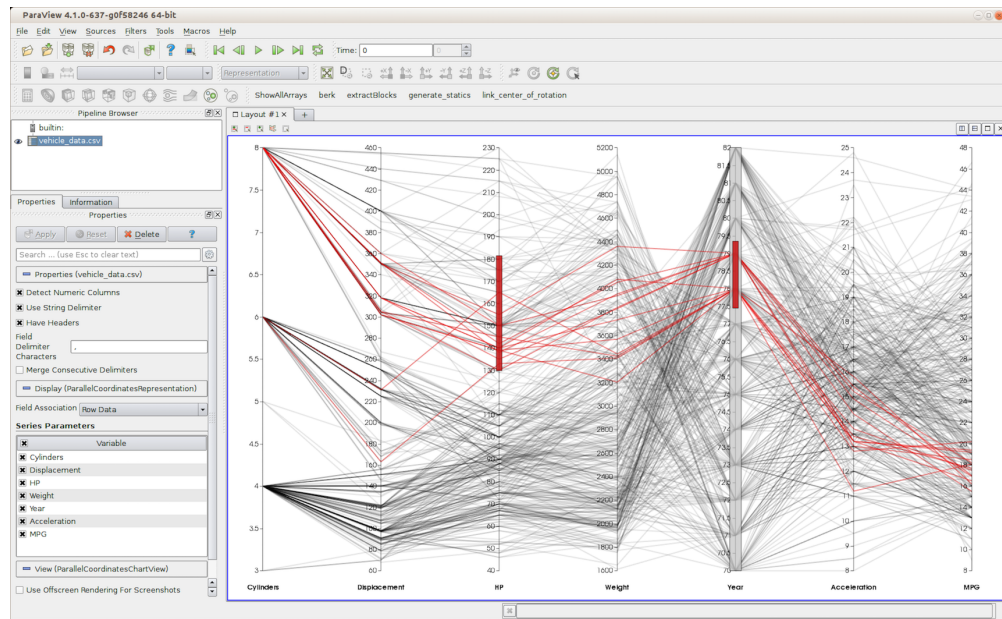


Fig. 1.48: paraview using Parallel Coordinates View to plot correlations between columns in a table.

Like Plot Matrix View, Parallel Coordinates View is also used to visualize correlations between data arrays.

One of the main features of this view is the ability to select specific data in order to analyse the factors influencing the data. e.g., with a table of three variables, one being the “output” variable the other two being the potential factor influencing the first, selecting only the output will enable you to see if none, one, or both of the factors are actually influencing the output.

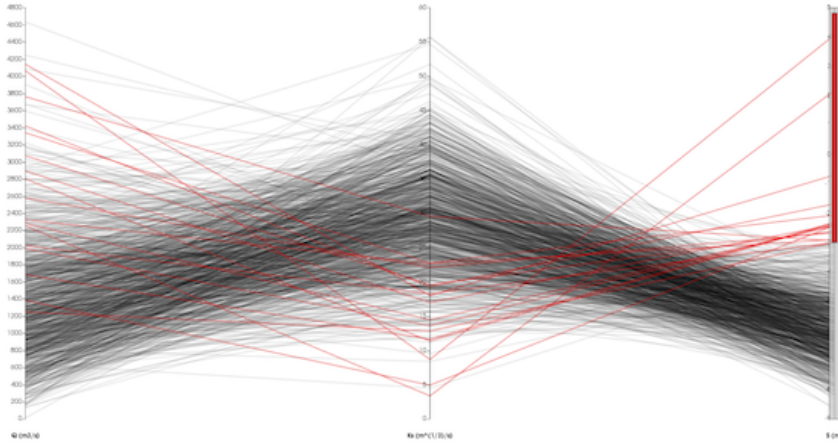


Fig. 1.49: High “S” data point are influenced more by low “Ks” than high “Q”

1.4.10 Spreadsheet View

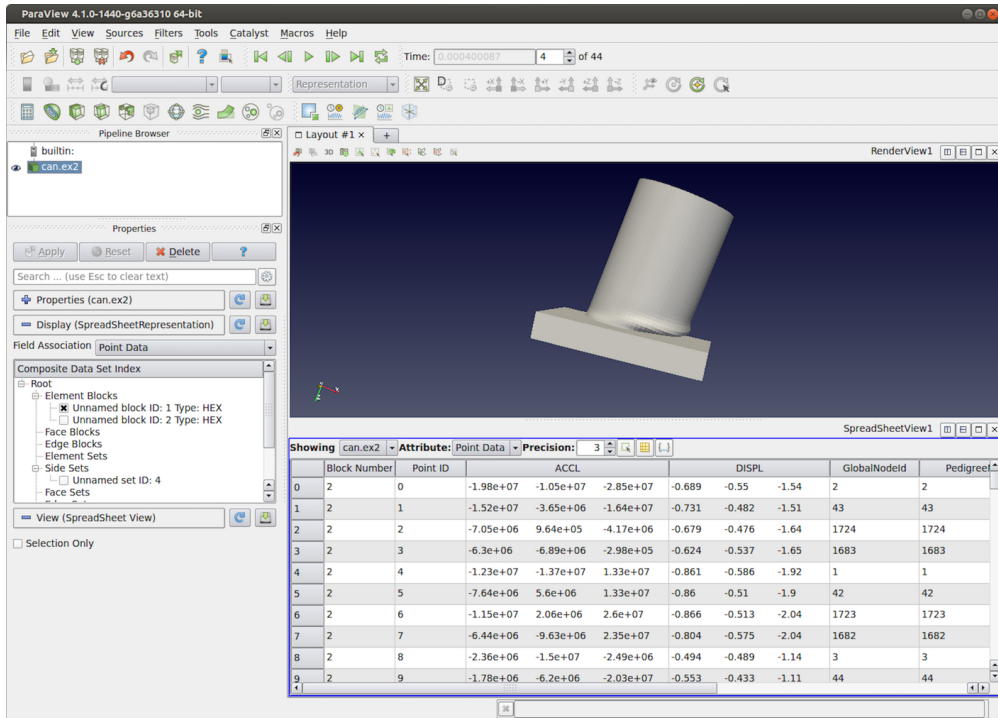


Fig. 1.50: paraview using SpreadSheet View to plot raw data values for the *can.ex2* dataset.

SpreadSheet View is used to inspect raw data values in a tabular form. Unlike most other views, this view is primarily intended to be used in the paraview user interface and, hence, is not available in pvython.

To use this view, simply create this view and show the dataset produced by any pipeline module in it by using the Pipeline Browser . SpreadSheet View can only show one dataset at a time. Therefore, showing a new dataset will automatically hide the previously shown dataset.

The view's toolbar provides quick access to several of this view's capabilities. Use the Showing widget on the view toolbar to view as well as to change the dataset being shown. The Attribute field allows you to pick which types of elements to show, e.g., Cell Data , Point Data , Field Data , etc. Precision can be utilized to change the precision used when displaying floating point numbers. The button enables you to select columns to show. Click on the button to get a popup menu in which you check/uncheck the columns to show/hide. If showing Cell Data , the button, when checked, enables you to see the point ids that form each of the cells.

Section 1.6.1 discusses how selections can be made in views to select elements of interest. Use the button to make the view show only selected elements. Now, as you make selections in other views, this SpreadSheet View will update to only show the values of the selected elements (as long as the dataset selected in are indeed being shown in the view).

Did you know?

Some filters and readers produce vtkTable, and they are automatically displayed in a spreadsheet view. Thus, one can very easily read the contents of a .csv file with ParaView.

1.4.11 Slice View

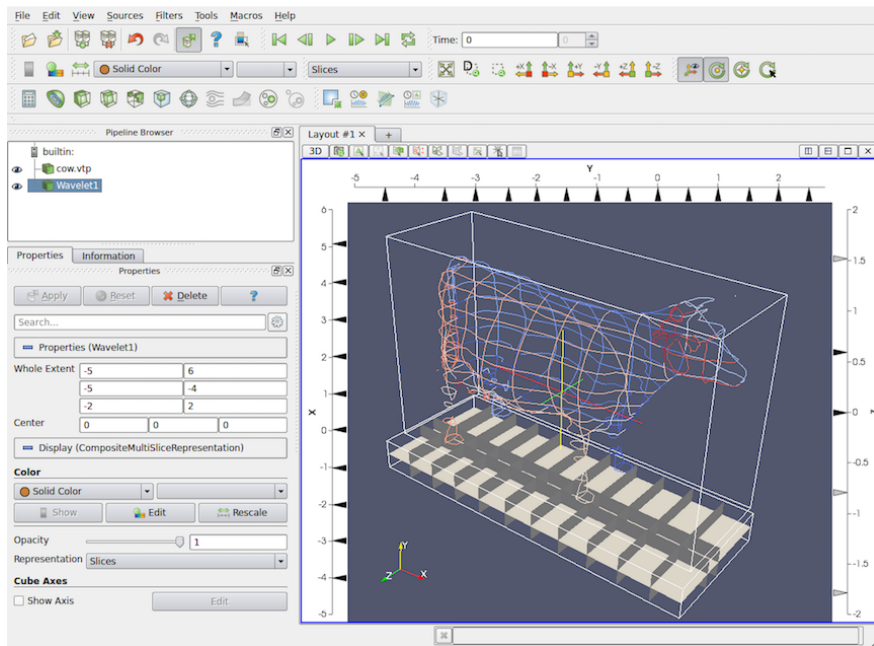


Fig. 1.51: Slice View can be used to show orthogonal slices from datasets.

Slice View is special type of Render View that can be used to view orthogonal slices from any dataset. Any dataset shown in the view will be sliced in axis-aligned slices based on the locations specified on the view. The slice locations along the three orthogonal axis planes can be specified by using the frame decoration around the view window.

Interactions

Since this view is a type of Render View, the camera interactions are same as that of Render View. Additionally, you can interact with the frame decoration to manipulate slice locations along the three axis planes.

- Double-click the left mouse button in the region between the axis border and the view to add a new slice.
- You can click-and-drag a marker to move the slice location.
- To remove a slice, double-click with the left mouse button on the marker corresponding to that slice.
- To toggle visibility of the slice, you can right-click on the marker.

Slice View in pvcppython

```
# To create a slice view in use:
>>> view = CreateView("MultiSlice")

# Use properties on view to set/get the slice offsets.
>>> view.XSliceValues = [-10, 0, 10]
>>> print(view.XSliceValues)
[-10, 0, 10]

# Similar to XSliceValues, you have YSliceValues and ZSliceValues.
```

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```
>>> view.YSliceValues = [0]
>>> view.ZSliceValues = []
```

1.4.12 Python View

Some Python libraries, such as matplotlib, are widely used for making publication-quality plots of data. The Python View provides a way to display plots made in a Python script right within paraview.

The Python View has a single property, a Python script that generates the image to be displayed in the viewport. All the Python bindings for ParaView and VTK that are available in the Python scripting module are available from within this script, making it possible to plot any array from just about any dataset that can be loaded into ParaView. The Python script for the view is evaluated in a unique Python environment so that global variables defined in the script do not clobber global variables in other Python scripts (either in other instances of the Python View or in the Python interpreter). This environment is reset each time the script is evaluated, so data cannot be saved between evaluations.

The Python View requires that the Python script where the plotting occurs define two functions. In the first function, you request which arrays you would like to transfer to the client for rendering. At present, all rendering in this view takes place on the client, even in client-server mode. These arrays can be point data, cell data, field data, and table row data. This function runs only on data-server processes. It provides access to the underlying data object on the server so that you can query any aspect of the data using the Python-wrapped parts of VTK and ParaView.

The second function is where you put Python plotting or rendering commands. This function runs only on the ParaView client. It has access to the complete data object gathered from the data server nodes, but only has access to the arrays requested in the first function. This function will typically set up objects from a plotting library, convert data from VTK to a form that can be passed to the plotting library, plot the data, and convert the plot to an image (a vtkImageData object) that can be displayed in the viewport.

Selecting data arrays to plot

All the rendering in the Python View occurs in the client, so the client must get the data from the server. Because the dataset residing on the ParaView server may be large, transferring all the data to the client may not be possible or practical. For that reason, we have provided a mechanism to select which data arrays in a data object on the server to transfer to the client. The overall structure of the data object, however, (including cell connectivity, point positions, and hierarchical block structure) is always transferred to the client. By default, no data arrays are selected for transfer from the server.

The Python script for the view must define a function called `setup_data(view)`. The `view` argument is the VTK object for the Python View. The current datasets loaded into ParaView may be accessed through the `view` object.

Here's an example of this function that was used to generate the image in Fig. 1.52:

```
def setup_data():
    # Iterate over visible data objects
    for i in xrange(view.GetNumberOfVisibleDataObjects()):
        # You need to use GetVisibleDataObjectForSetup(i)
        # in setup_data to access the data object.
        dataObject = view.GetVisibleDataObjectForSetup(i)

        # The data object has the same data type and structure
        # as the data object that sits on the server. You can
        # query the size of the data, for instance, or do anything
        # else you can do through the Python wrapping.
        print('Memory size: {} kilobytes'.format(dataObject.GetActualMemorySize()))
```

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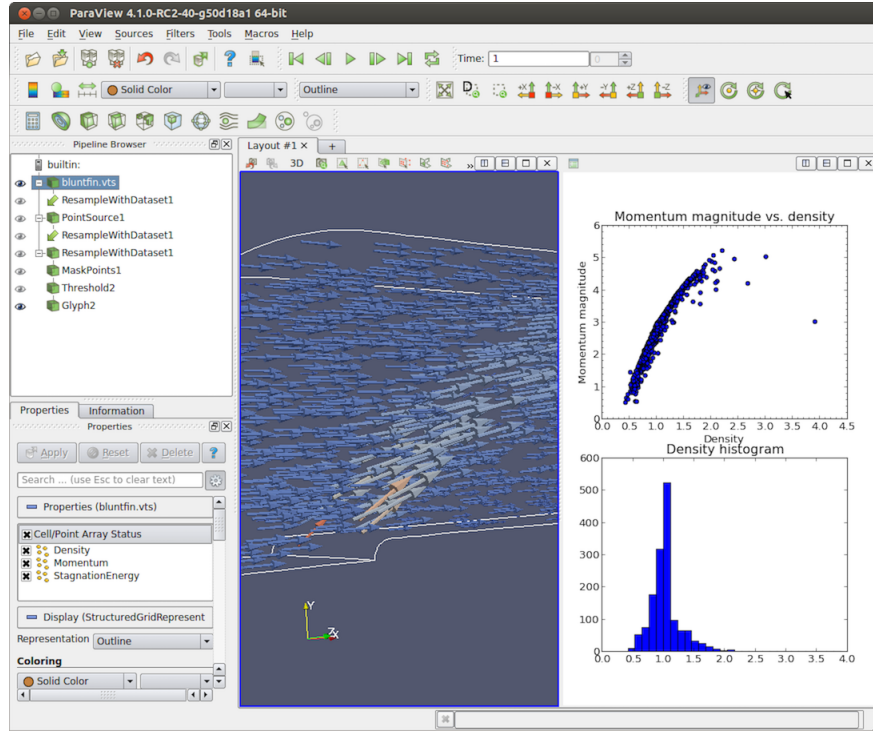


Fig. 1.52: paraview using Python View with matplotlib to display a scatterplot of momentum magnitude versus density (upper right) and a histogram of density (lower right) in the bluntfin.vts dataset.

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```
# Clean up from previous calls here. We want to unset
# any of the arrays requested in previous calls to this function.
view.DisableAllAttributeArrays()

# By default, no arrays will be passed to the client.
# You need to explicitly request the arrays you want.
# Here, we'll request the Density point data array
view.SetAttributeArrayStatus(i, vtkDataObject.POINT, "Density", 1)
view.SetAttributeArrayStatus(i, vtkDataObject.POINT, "Momentum", 1)

# Other attribute arrays can be set similarly
view.SetAttributeArrayStatus(i, vtkDataObject.FIELD, "fieldData", 1)
```

The `vtkPythonView` class passed in as the `view` argument to `setup_data(view)` defines several methods useful for specifying which data arrays to copy:

- `GetNumberOfVisibleDataObjects()` - This returns the number of visible data objects in the view. If an object is not visible, it should not show up in the rendering, so all the methods provided by the view deal only with visible objects.
- `GetVisibleDataObjectForSetup(visibleObjectIndex)` - This returns the `visibleObjectIndex`'th visible data object in the view. (The data object will have an open eye next to it in the pipeline browser.)
- `GetNumberOfAttributeArrays(visibleObjectIndex, attributeType)` - This returns the number of attribute arrays for the `visibleObjectIndex`'th visible object and the given `attributeType` (e.g., `vtkDataObject.POINT`, `vtkDataObject.CELL`, etc.).

- `GetAttributeArrayName(visibleObjectIndex, attributeType, arrayIndex)` - This returns the name of the array of the given attribute type at the given array index for the `visibleObjectIndex`'th object.
- `SetAttributeArrayStatus(visibleObjectIndex, vtkDataObject.POINT, "Density", 1)` - This sets the array status of an attribute array. The first argument is the visible object index, the second object is the attribute association of the array, the third argument is the name of the array, and the last argument specifies if the array is to be copied (1) or not (0).
- `GetAttributeArrayStatus(visibleObjectIndex, vtkDataObject.POINT, "Density")` - This retrieves the array status for the object with the given visible index with a given attribute association (second argument) and a name (last argument).
- `EnableAllAttributeArrays()` - This sets all arrays to be copied.
- `DisableAllAttributeArrays()` - This sets all arrays to not be copied.

The methods `GetNumberOfVisibleDataObjects()` , `GetVisibleDataObjectForSetup(...)` , `GetNumberOfAttributeArrays(...)` , and `GetAttributeArrayName(...)` are all convenient methods for obtaining information about visible data objects in the view that could otherwise be accessed with existing view and representation methods. The last four methods are valid only from within the `setup_data(view)` function.

Plotting data in Python

After the `setup_data(view)` function has been called, **ParaView** will transfer the data object and selected arrays to the client. When that is done, it will call the `render(view, width, height)` function you have defined in your script.

The `view` argument to the `render(view, width, height)` function is the `vtkPythonView` object on the client. The `width` and `height` arguments are the width and height of the viewport, respectively. The `render(view, width, height)` function uses the data available through the view, along with the width and height, to generate a `vtkImageData` object that will be displayed in the viewport. This `vtkImageData` object must be returned from the `render(view, width, height)` function. If no `vtkImageData` is returned, the viewport will be black. If the size of the image does not match the size of the viewport, the image will be stretched to fit the viewport.

Putting it all together, here is a simple example that generates a solid red image to display in the viewport.

```
def render(view, width, height):
    from paraview.vtk import vtkImageData
    image = vtkImageData()
    image.SetDimensions(width, height, 1)
    from paraview.numeric import VTK_UNSIGNED_CHAR
    image.AllocateScalars(VTK_UNSIGNED_CHAR, 4)
    pixel_array = image.GetPointData().GetArray(0)
    pixel_array.FillComponent(0, 255.0)
    pixel_array.FillComponent(1, 0.0)
    pixel_array.FillComponent(2, 0.0)
    pixel_array.FillComponent(3, 0.0)

    return image
```

This example does not produce an interesting visualization, but serves as a minimal example of how the `render(view, width, height)` function should be implemented. Typically, we expect that the Python plotting library you use has some utilities to expose the generated plot image pixel data. You need to copy that pixel data to the `vtkImageData` object returned by the `render(view, width, height)` function. Exactly how you do this is up to you, but **ParaView** comes with some utilities to make this task easier for matplotlib.

Set up a matplotlib Figure

The Python View comes with a Python module, called `python_view`, that has some utility functions you can use. To import it, use:

```
from paraview import python_view
```

This module has a function, called `matplotlib_figure(view, width, height)`, that returns a `matplotlib.Figure` given `width` and `height` arguments. This figure can be used with `matplotlib` plotting commands to plot data as in the following:

```
def render(view, width, height):
    figure = python_view.matplotlib_figure(width, height)

    ax = figure.add_subplot(1,1,1)
    ax.minorticks_on()
    ax.set_title('Plot title')
    ax.set_xlabel('X label')
    ax.set_ylabel('Y label')

    # Process only the first visible object in the pipeline browser
    dataObject = view.GetVisibleDataObjectForRendering(0)

    x = dataObject.GetPointData().GetArray('X')

    # Convert VTK data array to numpy array for plotting
    from paraview.numpy_support import vtk_to_numpy
    np_x = vtk_to_numpy(x)

    ax.hist(np_x, bins=10)

    return python_view.figure_to_image(figure)
```

This definition of the `render(view, width, height)` function creates a histogram of a point data array named `X` from the first visible object in the pipeline browser. Note the conversion function, `python_view.figure_to_image(figure)`, in the last line. This converts the `matplotlib Figure` object created with `python_view.matplotlib_figure(width, height)` into a `vtkImageData` object suitable for display in the viewport.

1.4.13 Comparative Views

Comparative Views , including Render View (Comparative) , Line Chart View (Comparative) , and Bar Chart View (Comparative) , are used for generating comparative visualization from parameter studies. We will cover these views in Chapter [Section 2.3](#).

1.5 Filtering Data

Visualization can be characterized as a process of transforming raw data produced from experiments or simulations until it takes a form in which it can be interpreted and analysed. The visualization pipeline introduced in [Section 1.1.2](#) formalizes this concept as a data flow paradigm where a pipeline is set up of sources, filters, and sinks (collectively called pipeline modules or algorithms). Data flows through this pipeline, being transformed at each node until it is in a form where it can be consumed by the sinks. In previous chapters, we saw how to ingest data into **ParaView** ([Section 1.2](#)) and how to display it in views ([Section 1.4](#)). If the data ingested into **ParaView** already has all the relevant attribute data, and it is in the form that can be directly represented in one the existing views, then that is all you would need. The true power of the visualization process, however, comes from leveraging the various visualization techniques such as slicing, contouring, clipping, etc., which are available as filters. In this chapter, we look at constructing pipelines to transform data using such filters.

1.5.1 Understanding filters

In **ParaView**, filters are pipeline modules or algorithms that have inputs and outputs. They take in data on their inputs and produce transformed data or results on their outputs. A filter can have multiple input and output ports. The number of input and output ports on a filter is fixed. Each input port accepts input data for a specific purpose or role within the filter. (E.g., the Resample With Dataset filter has two input ports. The one called Input is the input port through which the dataset providing the attributes to interpolate is ingested. The other, called Source , is the input port through which the dataset used as the mesh on which to re-sample is accepted.)

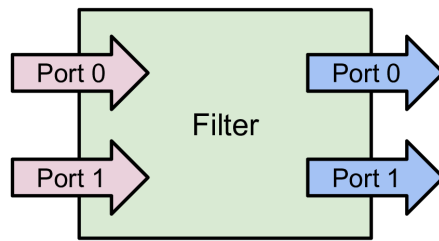


Fig. 1.53: A filter is a pipeline module with inputs and outputs. Data enters a filter through the inputs. The filter transforms the data and produces the resulting data on its outputs. A filter can have one or more input and output ports. Each input port can optionally accept multiple input connections.

An input port itself can optionally accept multiple input connections, e.g., the Append Datasets filter, which appends multiple datasets to create a single dataset only has one input port (named Input). However, that port can accept multiple connections for each of the datasets to be appended . Filters define whether a particular input port can accept one or many input connections.

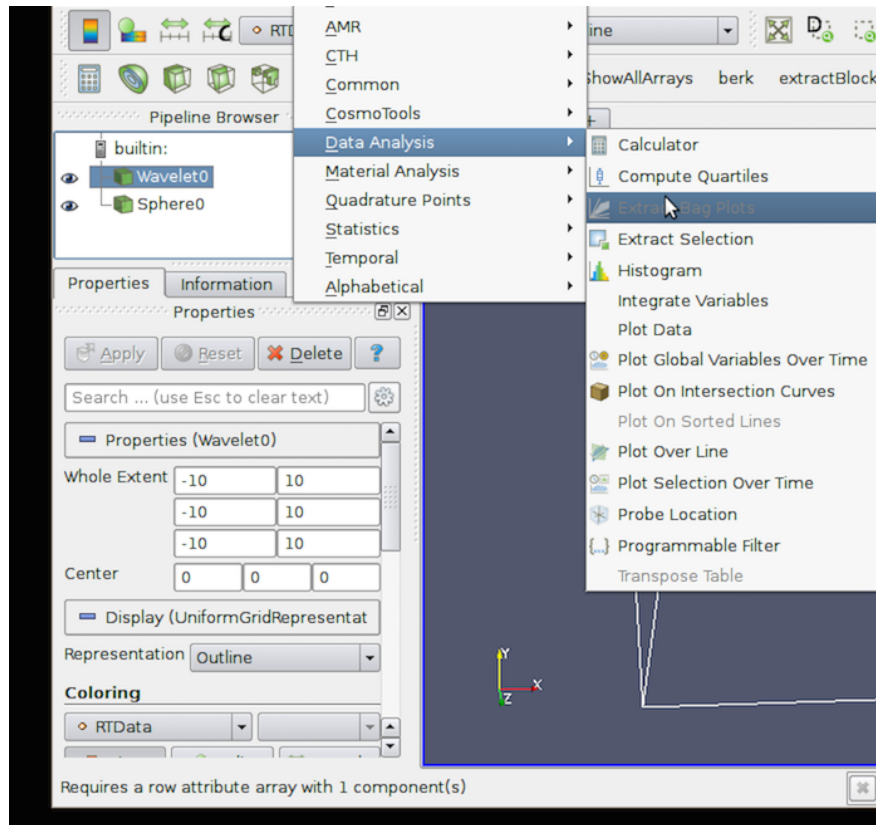
Similar to readers, the properties on the filter allow you to control the filtering algorithm. The properties available depend on the filter itself.

1.5.2 Creating filters in paraview

All available filters in `paraview` are listed under the *Filters* menu. These are organized in various categories. To create a filter to transform the data produced by a source or a reader, you select the source in the `Pipeline Browser` to make it active, and then click on the corresponding menu item in the *Filters* menu. If a menu item is disabled, it implies that the active source does not produce data that can be transformed by this filter.

Did you know?

If a menu item in the *Filters* menu is disabled, it implies that the active source(s) is not producing data of the expected type or the characteristics needed by the filter. On Windows and Linux machines, if you hover over the disabled menu item, the status bar will show the reason why the filter is not available.



Multiple input connections

When you create a filter, the active source is connected to the first input port of the filter. Filters like `Append Datasets` can take multiple input connections on that input port. In such a case, to pass multiple pipeline modules as connections on a single input port of a filter, select all the relevant pipeline modules in the `Pipeline Browser`. You can select multiple items by using the `CTRL` (or `Cmd`) and key modifiers. When multiple pipeline modules are selected, only the filters that accept multiple connections on their input ports will be enabled in the *Filters* menu.

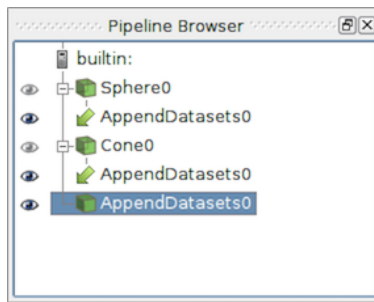


Fig. 1.54: The Pipeline Browser showing a pipeline with multiple input connections. The Append Datasets filter has two input connections on its only input port, `Sphere0` and `Cone0`.

Multiple input ports

Most filters have just one input port. Hence, as soon as you click on the filter name in the *Filters* menu, it will create a new filter instance and that will show up in the Pipeline Browser. Certain filters, such as `Resample With Dataset`, have multiple inputs that must be set up before the filter can be created. In such a case, when you click on the filter name, the `Change Input Dialog` will pop up, as seen in Fig. 1.55. This dialog allows you to select the pipeline modules to be connected to each of the input ports. The active source(s) is connected by default to the first input port. You are free to change those as well.

Changing input connections

paraview allows you to change the inputs to a filter after the filter has been created. To change inputs to a filter, right-click on the filter in the Pipeline Browser to get the context menu, and then select `Change Input...`. This will pop up the same `Change Input Dialog` as when creating a filter with multiple input ports. You can use this dialog to set new inputs for this filter.

Did you know?

While the *Filters* menu is a handy way to create new filters, with the long list of filters available in **ParaView**, manually finding a particular filter in this menu can be very challenging. To make it easier, **ParaView** incorporates a quick launch mechanism. When you want to create a new filter (or a source), simply type **CTRL + Space** or **Alt + Space**. This will pop up the quick-launch dialog. Now, start typing the name of the filter you want. As you type, the dialog will update to show the filters and sources that match the typed text. You can use the arrow keys to navigate and use the **Enter** key to create the selected filter (or source). Press while pressing **Enter** to quickly apply the filter on creation, equivalent to creating the filter and then clicking the **Apply** button. Note that filters may be disabled, as was the case in the *Filters* menu but by default the selected item will be the first enabled filter. You can use **Esc** to clear the text you have typed so far. Hit the **Esc** a second time, and the dialog will close without creating any new filter.

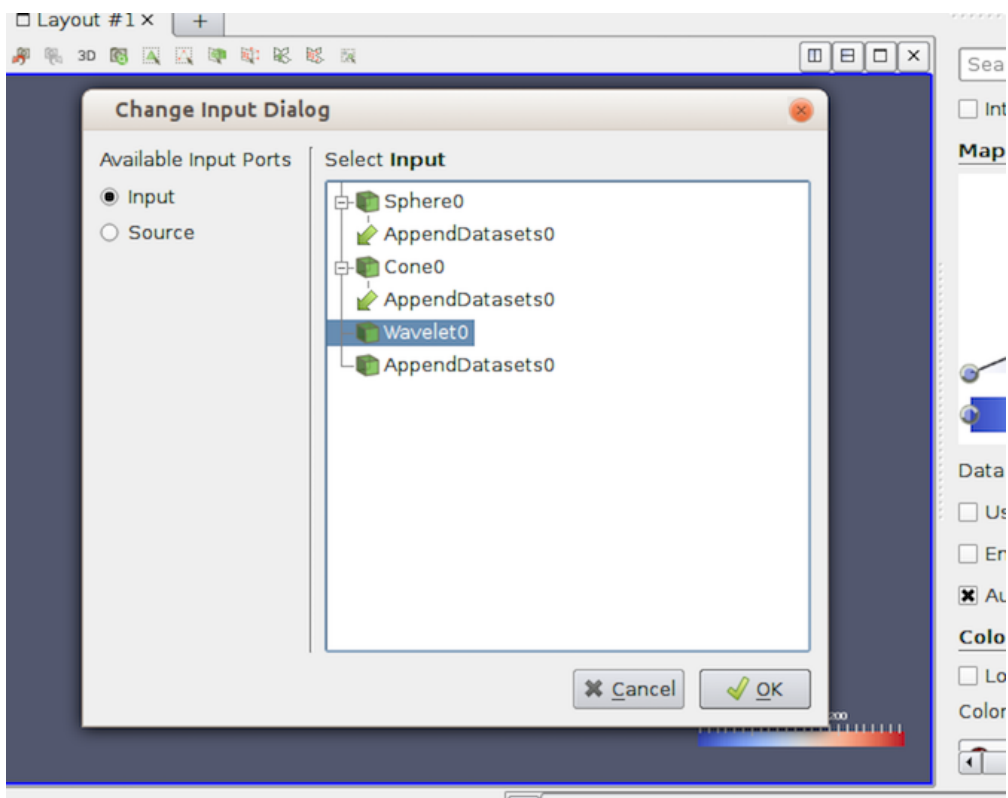


Fig. 1.55: The `Change Input Dialog` is shown to allow you to pick inputs for each of the input ports for a filter with multiple input ports. To use this dialog, first select the `Input Port` you want to edit on the left side, and select the pipeline module(s) that are to be connected to this input port. Repeat the step for the other input port(s). If an input port can accept multiple input connections, you can select multiple modules, just like in the `Pipeline Browser`.

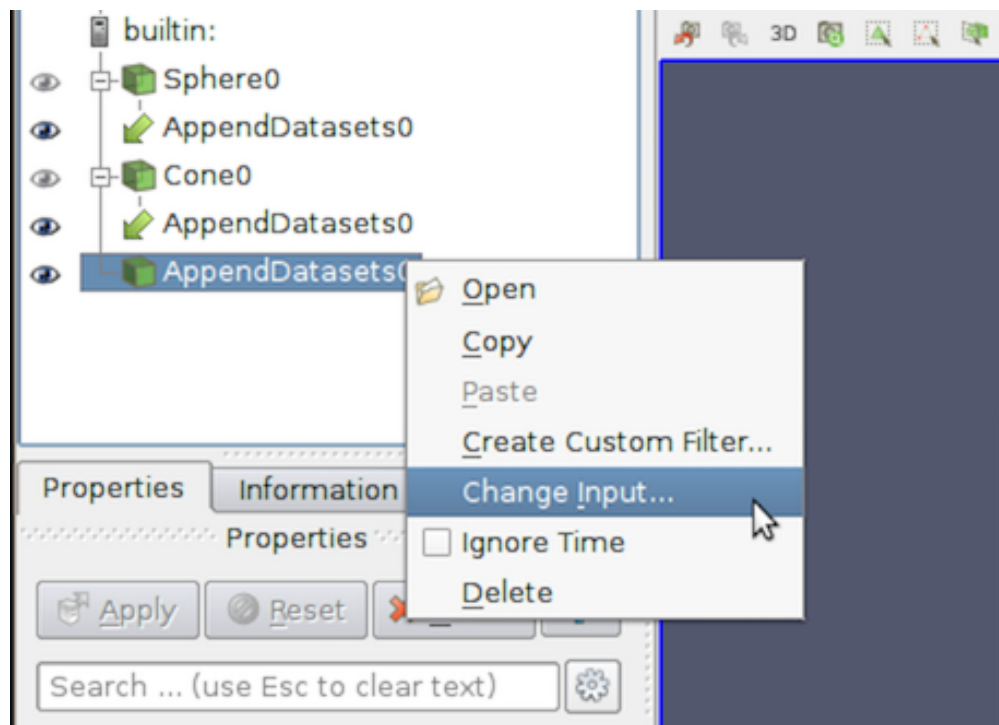
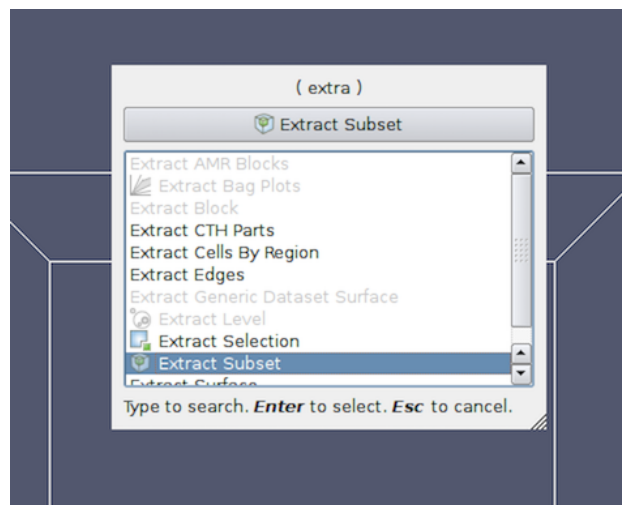


Fig. 1.56: The context menu in the Pipeline Browser showing the option to change inputs for a filter.



1.5.3 Creating filters in pypython

To create a filter in pypython, you simply create the object by using its name as a constructor function.

```
>>> from paraview.simple import *
...
>>> filter = Shrink()
```

Similar to paraview, the filter will use the active source(s) as the input. Additionally, you can explicitly specify the input in the function arguments.

```
>>> reader = OpenDataFile(...)
...
>>> shrink = Shift(Input=reader)
```

Multiple input connections

To setup multiple input connections, you can specify the connections as follows:

```
>>> sphere = Sphere()
>>> cone = Cone()

# Simply pass the sources as a list to the constructor function.
>>> appendDatasets = AppendDatasets(Input=[sphere, cone])
>>> print(appendDatasets.Input)
[<paraview.servermanager.Sphere object at 0x6d75f90>, <paraview.servermanager.Cone_
object at 0x6d75c50>]
```

Multiple input ports

Setting up connections to multiple input ports is similar to the multiple input connections, except that you need to ensure that you name the input ports properly.

```
>>> sphere = Sphere()
>>> wavelet = Wavelet()

>>> resampleWithDataSet = ResampleWithDataSet(Input=sphere, Source=wavelet)
```

Changing input connections

Changing inputs in Python is as simple as setting any other property on the filter.

```
# For filter with single input connection
>>> shrink.Input = cone

# for filters with multiple input connects
>>> appendDatasets.Input = [reader, cone]

# to add a new input.
>>> appendDatasets.Input.append(sphere)
```

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```
# to change multiple ports
>>> resampleWithDataSet.Input = wavelet2
>>> resampleWithDataSet.Source = cone
```

1.5.4 Changing filter properties in paraview

Filters provide properties that you can change to control the processing algorithm employed by the filter. Changing and viewing properties on filters is the same as with any other pipeline module, including readers and sources. You can view and change these properties, when available, using the **Properties** panel. Section 2.1 covers how to effectively use the **Properties** panel. Since this panel only shows the properties present on the *active source*, you must ensure that the filter you are interested in is active. To make the filter active, use the **Pipeline Browser** to click on the filter and select it.

1.5.5 Changing filter properties in pvvpython

With pvvpython, the available properties are accessible as properties on the filter object, and you can get or set their values by name (similar to changing the input connections (Section 1.5.3)).

```
# You can save the object reference when it's created.
>>> shrink = Shrink()

# Or you can get access to the active source.
>>> Shrink() # <-- this will make the Shrink the active source.
>>> shrink = GetActiveSource()

# To figure out available properties, you can always use help.
>>> help(shrink)
Help on Shrink in module paraview.servermanager object:

class Shrink(SourceProxy)
|   The Shrink filter
|   causes the individual cells of a dataset to break apart
|   from each other by moving each cell's points toward the
|   centroid of the cell. (The centroid of a cell is the
|   average position of its points.) This filter operates on
|   any type of dataset and produces unstructured grid
|   output.
|
|   Data descriptors defined here:
|
|   Input
|       This property specifies the input to the Shrink
|       filter.
|
|   ShrinkFactor
|       The value of this property determines how far the points
|       will move. A value of 0 positions the points at the centroid of the
|       cell; a value of 1 leaves them at their original
|       positions.
```

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```
....  
  
# To get the current value of a property:  
>>> sf = shrink.ShrinkFactor  
>>> print sf  
0.5  
  
# To set the value  
>>> shrink.ShrinkFactor = 0.75
```

In the rest of this chapter, we will discuss some of the commonly used filters in detail. They are grouped under categories based on the type of operation that they perform.

1.5.6 Filters for sub-setting data

These filters are used for extracting subsets from an input dataset. How this subset is defined and how it is extracted depends on the type of the filter.

Clip

Clip is used to clip any dataset using either an implicit function (such as a plane, sphere, or a box) or using values of a scalar data array in the input dataset. A scalar array is a point or cell attribute array with a single component. Clipping involves iterating over all cells in the input dataset and then removing those cells that are considered *outside* of the space defined by the implicit function or that have an attribute values less than the selected value. For cells that straddle the clipping surface, these are *clipped* to pass through the part of the cell that is truly inside the specified implicit function (or greater than the scalar value).

This filter converts any dataset into an unstructured grid (Section 1.3.1) or a multi-block of unstructured grids (Section 1.3.1) in the case of composite datasets.

Clip in paraview

To create the Clip filter, you can use the *Filters > Common* or the *Filters > Alphabetical* menu. This filter is also accessible from the Common filters toolbar. You can click the  button to create this filter.

On the Properties panel, you will see the available properties for this filter. One of the first things that you should select is the Clip Type . Clip Type is used to specify the type of implicit function to use for the clipping operations. The available options include Plane , Box , Sphere , and Scalar . Selecting any one of these options will update the panel to show properties that are used to define the implicit function, e.g., the Origin and the Normal for the Plane or the Center and the Radius for the Sphere . If you select Scalar , the panel will let you pick the data array and the value with which to clip. Remember, cells with the data value greater than or equal to the selected value are considered *in* and are passed through the filter.

Did you know?

When clipping with implicit functions, **ParaView** renders widgets in the active view that you can use to interactively control the implicit function, called 3D widgets . As you interact with the 3D widget, the panel will update to reflect the current values. The 3D widget is considered as an aid and not as a part of the actual visualization scene. Thus, if you change the active source and the Properties panel navigates away from this filter, the 3D widget will automatically be hidden.

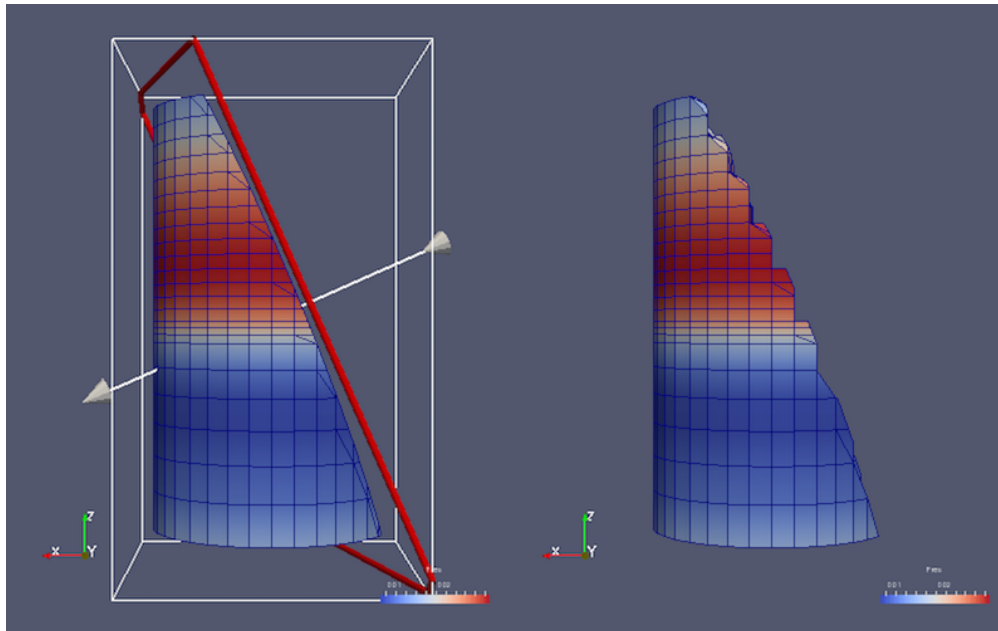


Fig. 1.57: Comparison between results produced by the `Clip` filter with `Crinkle Clip` unchecked (left) and checked (right) when clipping with an implicit plane. The image on the left also shows the 3D widget used to interactively place the implicit plane for the clipping operation.

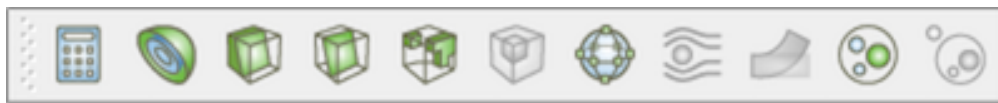


Fig. 1.58: The Common filters toolbar in paraview for quick access to the commonly used filters.

The `Inside Out` option can be used to invert the behavior of this filter. Basically, it flips the notion of what is considered inside and outside of the given clipping space.

Check `Crinkle Clip` if you don't want this filter to truly clip cells on the boundary, but want to preserve the input cell structure and to pass the entire cell on through the boundary (Fig. 1.57). This option is not available when clipping by `Scalar`.

Clip in pvvpython

This following script demonstrates various aspects of using the `Clip` filter in pvvpython.

```
# Create the Clip filter.
>>> clip = Clip(Input=...)

# Specify a 'ClipType' to use.
>>> clip.ClipType = 'Plane'

# You can also use the SetProperties API instead.
>>> SetProperties(clip, ClipType='Plane')

>>> print(clip.GetProperty('ClipType').GetAvailable())
['Plane', 'Box', 'Sphere', 'Scalar']

# To set the plane origin and normal
>>> clip.ClipType.Origin = [0, 0, 0]
>>> clip.ClipType.Normal = [1, 0, 0]

# If you want to change to Sphere and set center and
# radius, you can do the following.
>>> clip.ClipType = 'Sphere'
>>> clip.ClipType.Center = [0, 0, 0]
>>> clip.ClipType.Radius = 12

# Using SetProperties API, the same looks like
>>> SetProperties(clip, ClipType='Sphere')
>>> SetProperties(clip.ClipType, Center=[0, 0, 0],
                  Radius = 12)

# To set Crinkle clipping.
>>> clip.Crinkleclip = 1

# For clipping with scalar, you pick the scalar array
# and then the value as follows:
>>> clip.ClipType = 'Scalar'
>>> clip.Scalars = ('POINTS', 'Temp')
>>> clip.Value = 100
```

```
# As always, to get the list of available properties on
# the clip filter, use help()
>>> help(clip)
Help on Clip in module paraview.servermanager object:

class Clip(SourceProxy)
```

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```

| The Clip filter
| cuts away a portion of the input dataset using an
| implicit plane. This filter operates on all types of data
| sets, and it returns unstructured grid data on
| output.

-----
| Data descriptors defined here:

| ClipType
|     This property specifies the parameters of the clip
|     function (an implicit plane) used to clip the dataset.

| Crinkleclip
|     This parameter controls whether to extract entire cells
|     in the given region or clip those cells so all of the output one stay
|     only inside that region.

| Input
|     This property specifies the dataset on which the Clip
|     filter will operate.

| InsideOut
|     If this property is set to 0, the clip filter will
|     return that portion of the dataset that lies within the clip function.
|     If set to 1, the portions of the dataset that lie outside the clip
|     function will be returned instead.

...
# To get help on a specific implicit function type, make it the active
# ClipType and then use help()
>>> clip.ClipType = 'Plane'
>>> help(clip.ClipType)
Help on Plane in module paraview.servermanager object:

class Plane(Proxy)
...

```

Common Errors

It is very easy to forget that clipping a structured dataset such as image data can dramatically increase the memory requirements, since this filter will convert the structured dataset into an unstructured grid due to the nature of the clipping operation itself. For structured dataset, think about using **Slice** or **Extract Subset** filters instead, whenever appropriate. Those are not entirely identical operations, but they are often sufficient.

Slice

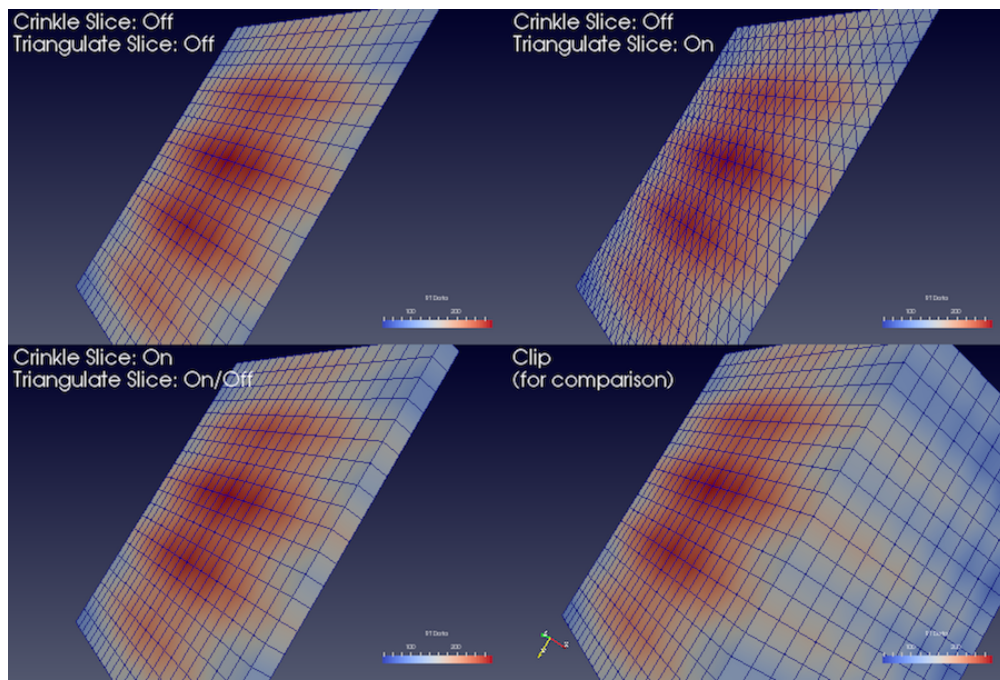


Fig. 1.59: Comparison between results produced by the `Slice` filter when slicing image data with an implicit plane with different options. The lower-left image shows the output produced by the `Clip` filter when clipping with the same implicit function, for contrast.

The `Slice` filter slices through the input dataset with an implicit function such as a plane, a sphere, or a box. Since this filter returns data elements along the implicit function boundary, this is a dimensionality reducing filter (except when crinkle slicing is enabled), i.e., if the input dataset has 3D elements like tetrahedrons or hexahedrons, the output will have 2D elements, line triangles, and quads, if any. While slicing through a dataset with 2D elements, the result will be lines.

The properties available on this filter, as well as the way of setting this filter up, is very similar to the `Clip` filter with a few notable differences. What remains similar is the set up of the implicit function – you have similar choices: `Plane`, `Box`, `Sphere`, and `Cylinder`, as well as the option to toggle `Crinkle slice` (i.e., to avoid cutting through cells, pass complete cells from the input dataset that intersects the implicit function).

What is different includes the lack of slicing by `Scalar` (for that, you can use the `Contour` filter) and a new option, `Triangulate the slice`. Fig. 1.59 shows the difference in the generated meshes when various slice properties are changed.

The `Slice` filter is more versatile than the `Slice` representation. First, the `Slice` representation is available for image datasets only, whereas the `Slice` filter can be used on any type of 3D dataset. Second, the representation extracts a subset of the image consisting of a 2D slice oriented in the XY, YZ, or XZ planes at the image voxel locations while the plane used by the filter can be placed arbitrarily. Third, since the `Slice` representation always shows a flat object and lighting may interfere with interpretation of data values on the slice, lighting is not applied to the `Slice` representation. Lighting is applied, however, to results from the `Slice` filter. Lastly, the `Slice` representation may be faster than the filter to update and scrub through different slices because it does not need to compute the intersection of a plane with cells in the dataset.

In paraview, this filter can be created using the  button on the `Common` filters toolbar, besides the `Filters` menu.

Extract Subset

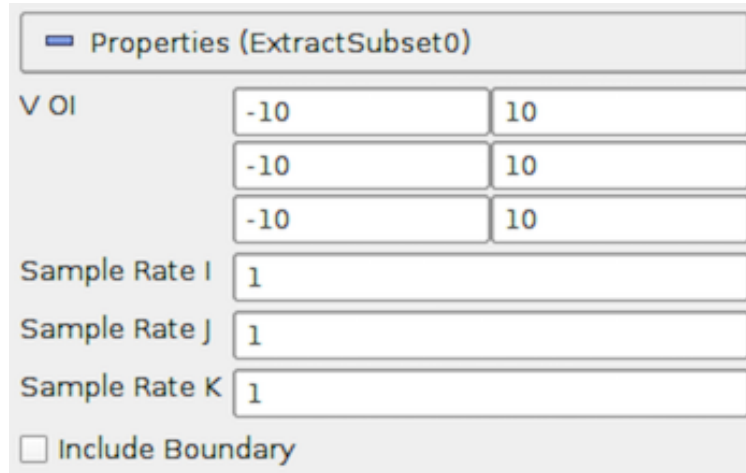


Fig. 1.60: The Properties panel for the `Extract Subset` filter showing all available properties (including the advanced properties).

For structured datasets such as image datasets (Section 1.3.1), rectilinear grids (Section 1.3.1), and curvilinear grids (Section 1.3.1), `Extract Subset` filter can be used to extract a region of interest or a subgrid. The region to extract is specified using structured coordinates, i.e., the i, j, k values. Whenever possible, this filter should be preferred over `Clip` or `Slice` for structured datasets, since it preserves the input data type. Besides extracting a subset, this filter can also be used to resample the dataset to a coarser resolution by specifying the sample rate along each of the structured dimensions.

Extract Subset in paraview

This is one of the filters available on the Common filters toolbar  To specify the region of interest, use the `VOI` property. The values are specified as min and max values for each of the structured dimensions (i, j, k) in each row. `Sample Rate I`, `Sample Rate J`, and `Sample Rate K` specify the sub-sampling rate. Set it to a value greater than one to sub-sample. `Include Boundary` is used to determine if the boundary slab should be included in the extracted result, if the sub-sampling rate along that dimension is greater than 1, and the boundary slab would otherwise have been skipped.

Threshold

The `Threshold` filter extracts cells of the input dataset with scalar values lying within the specified range, depending on the selected threshold method. This filter operates on either point-centered or cell-centered data. Any type of dataset can be used as input. The filter produces an unstructured grid output.

When thresholding with cell data, all cells that have scalars within the specified range will be passed through the filter. When thresholding with point data, cells with *all* points with scalar values within the range are passed through if `All Scalars` is checked; otherwise, cells with *any* point that passes the thresholding criteria are passed through.

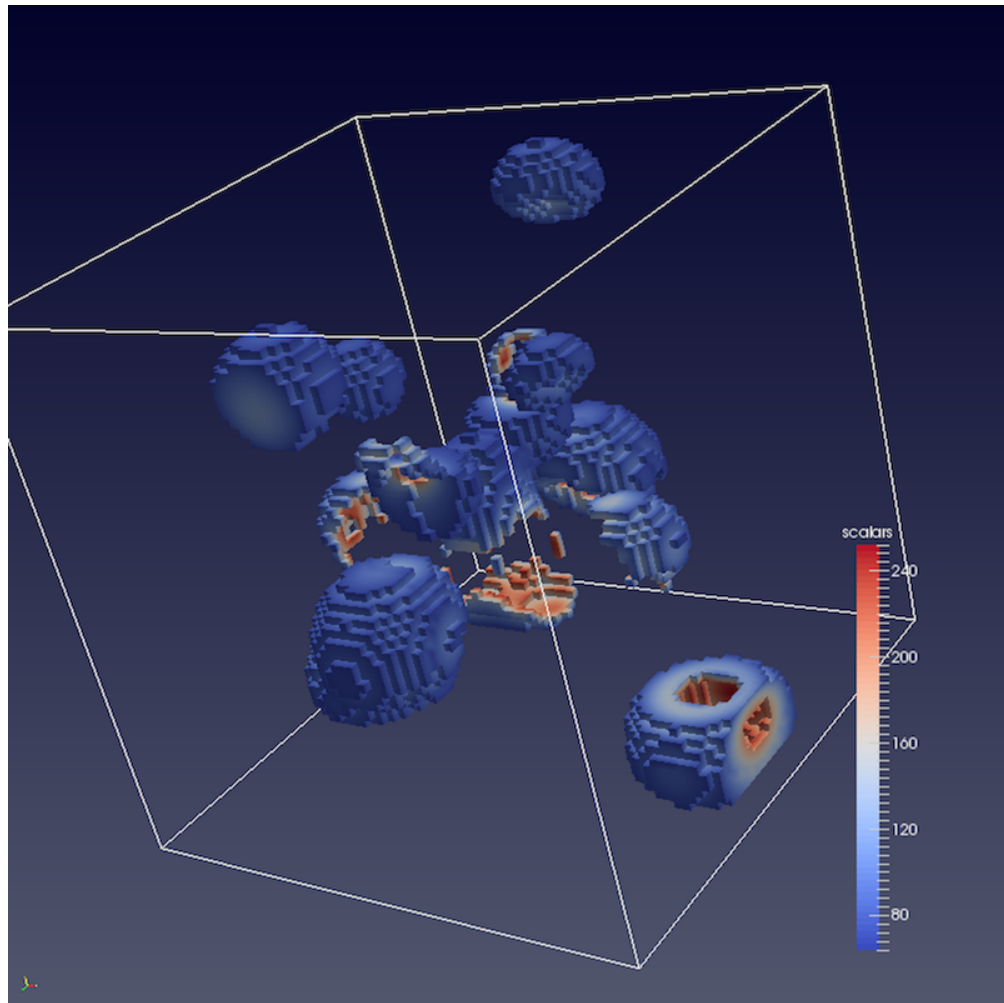


Fig. 1.61: Results from using the Threshold filter on the *iron_protein.vtk* dataset from **ParaView** data.

Threshold in paraview

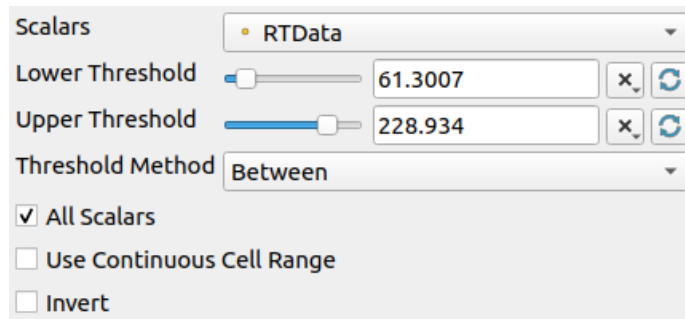


Fig. 1.62: The Properties panel for the Threshold filter.

This filter is represented as  on the Common filters toolbar. After selecting the Scalars with which to threshold from the combo-box, the Lower Threshold and Upper Threshold values can be modified to specify the range. If the range shown by the sliders is not sufficient, it is also possible to manually type the values in the input boxes. The values are deliberately not clamped to the current data range.

The threshold method can also be selected using the Threshold Method combo box:

- Between: Extracts cells with scalar values between the Lower Threshold and Upper Threshold.
- Below Lower Threshold: Extracts cells with scalar values smaller than the Lower Threshold.
- Above Upper Threshold: Extracts cells with scalar values larger than the Upper Threshold.

Threshold in pvvpython

```
# Create the filter. If Input is not specified, the active source will be
# used as the input.
>>> threshold = Threshold(Input=...)

# Here's how to select a scalar array.
>>> threshold.Scalars = ("POINTS", "scalars")

# The value is a tuple where the first value is the association: either "POINTS"
# or "CELLS", and the second value is the name of the selected array.
>>> print(threshold.Scalars)
['POINTS', 'scalars']

>>> print(threshold.Scalars.GetArrayName())
'scalars'

>>> print(threshold.Scalars.GetAssociation())
'POINTS'

# Different threshold methods are available and are set using one of the following:
>>> threshold.ThresholdMethod = "Between"                      # Uses both lower and upper
   ↵values
>>> threshold.ThresholdMethod = "Below Lower Threshold"    # Uses only lower value
>>> threshold.ThresholdMethod = "Above Upper Threshold"     # Uses only upper value
```

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```
# The adequate threshold values are then specified as:  
>>> threshold.LowerThreshold = 63.75  
>>> threshold.UpperThreshold = 252.45
```

To determine the types of arrays available in the input dataset, and their ranges, refer to the discussion on data information in [Section 1.3.3](#).

Iso Volume

The Iso Volume filter is similar to Threshold in that you use this to create an output dataset from an input where the cells that satisfy the specified range are scalar values. In fact, the filter is identical to Threshold when the cell data scalars are selected. For point data scalars, however, this filter acts similar to the Clip filters when clipping with scalars, in that cells are clipped along the iso-surface formed by the scalar range.

Extract Selection

Extract Selection is a general-purpose filter to extract selected elements from a dataset. There are several ways of making selections in ParaView. Once you have made the selection, this filter allows you to extract the selected elements as a new dataset for further processing. We will cover this filter in more detail when looking at selections in ParaView in [Section 1.6.6](#).

1.5.7 Filters for geometric manipulation

These filters are used to transform the geometry of the dataset without affecting its topology or its connectivity.

Transform

The Transform can be used to arbitrarily translate, rotate, and scale a dataset. The transformation is applied by scaling the dataset, rotating it, and then translating it based on the values specified.

As this is a geometric manipulation filter, this filter does not affect connectivity in the input dataset. While it tries to preserve the input dataset type, whenever possible, there are cases when the transformed dataset can no longer be represented in the same data type as the input. For example, with image data ([Section 1.3.1](#)) and rectilinear grids ([Section 1.3.1](#)) that are transformed by rotation, the output dataset can be non-axis aligned and, hence, cannot be represented as either data types. In such cases, the dataset is converted to a structured, or curvilinear, grid ([Section 1.3.1](#)). Since curvilinear grids are not as compact as the other two, the need to store the results in a more general data type implies a considerable increase in the memory footprint.

Transform in paraview

You can create a new Transform from the *Filters > Alphabetical* menu. Once created, you can set the transform as the translation, rotation, and scale to use utilizing the Properties panel. Similar to Clip, this filter also supports using a 3D widget to interactively set the transformation.

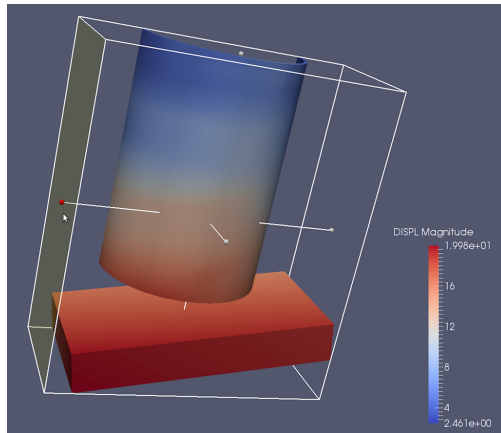


Fig. 1.63: The `Transform` filter showing the 3D widget that can be used to interactively set the transform.

Transform in pypython

```
# To create the filter(if Input is not specified, the active source will be
# used as the input).
>>> transform = Transform(Input=...)

# Set the transformation properties.
>>> transform.Transform.Scale = [1, 2, 1]
>>> transform.Transform.Translate = [100, 0, 0]
>>> transform.Transform.Rotate = [0, 0, 0]
```

Reflect

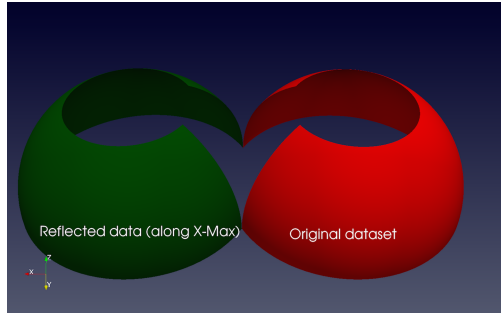


Fig. 1.64: The `Reflect` filter can be used to reflect a dataset along a specific axis plane.

`Reflect` can be used to reflect any dataset across an axis plane. You can pick the axis plane to be one of the planes formed by the bounding box of the dataset. For that, set `Plane` as `X Min`, `X Max`, `Y Min`, `Y Max`, `Z Min`, or `Z Max`. To reflect across an arbitrary axis plane, select `X`, `Y`, or `Z` for the `Plane` property, and then set the `Center` to the plane offset from the origin.

This filter reflects the input dataset and produces an unstructured grid (Section 1.3.1). Thus, the same caveats for `Clip` and `Threshold` filter apply here when dealing with structured datasets.

Warp By Vector

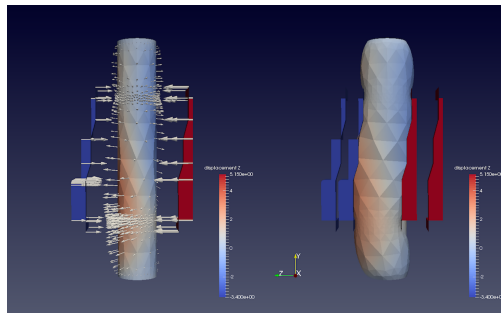


Fig. 1.65: The **Warp By Vector** filter can be used to displace points in original data shown on the left, using the *displacement* vectors (indicated by arrow glyphs Section 1.5.8) to produce the result shown on the right.

Warp By Vector can be used to displace point coordinates in an input mesh using vectors in the dataset itself. You select the vectors to use utilizing the **Vectors** property on the Properties panel. **Scale Factor** can be used to scale the displacement applied.

Warp By Scalar

Warp By Scalar is similar to **Warp By Vector** in the sense that it warps the input mesh. However, it does so using a scalar array in the input dataset. The direction of displacement can either be explicitly specified using the **Normal** property, or you can check **Use Normal** to use normals at the point locations.

1.5.8 Filters for sampling

These filters compute new datasets that represent some essential features from the datasets that they take as input.

Glyph

Glyph is used to place markers or glyphs at point locations in the input dataset. The glyphs can be oriented or scaled based on vector and scalar attributes on those points.

To create this filter in paraview, you can use the *Filters* menu, as well as the button on the Common filters toolbar. You first select the type of glyph using one of the options in **Glyph Type**. The choices include **Arrow**, **Sphere**, **Cylinder**, etc. Next, you select the point arrays to use as the **Orientation Array** (selecting **No orientation array** will result in the glyphs not being oriented). Similarly, you select a point array to serve as the **glyph Scale Array** (no scaling is performed if **No scale array** is chosen).

If the **Scale Array** is set to a vector array, the **Vector Scale Mode** property is available to select which properties of the vector should be used to transform each glyph. If **Scale by Magnitude** is chosen, then the glyph at a point will be scaled by the magnitude of the vector at that point. If **Scale by Components** is chosen, glyphs will be scaled separately in each dimension by the vector component in that dimension.

The **Scale Factor** is used to apply a constant scaling to all the glyphs, independent of the **Scale Array** and **Vector Scale Mode** properties. Choosing a good scale factor depends on several things including the bounds on the input dataset, the **Scale Array** and **Vector Scale Mode** selected, and the range for the array selected as the **Scale Array**. You can use the button next to the **Scale Factor** widget to have paraview pick a usually reasonable scale factor value based on the current dataset and scaling properties.

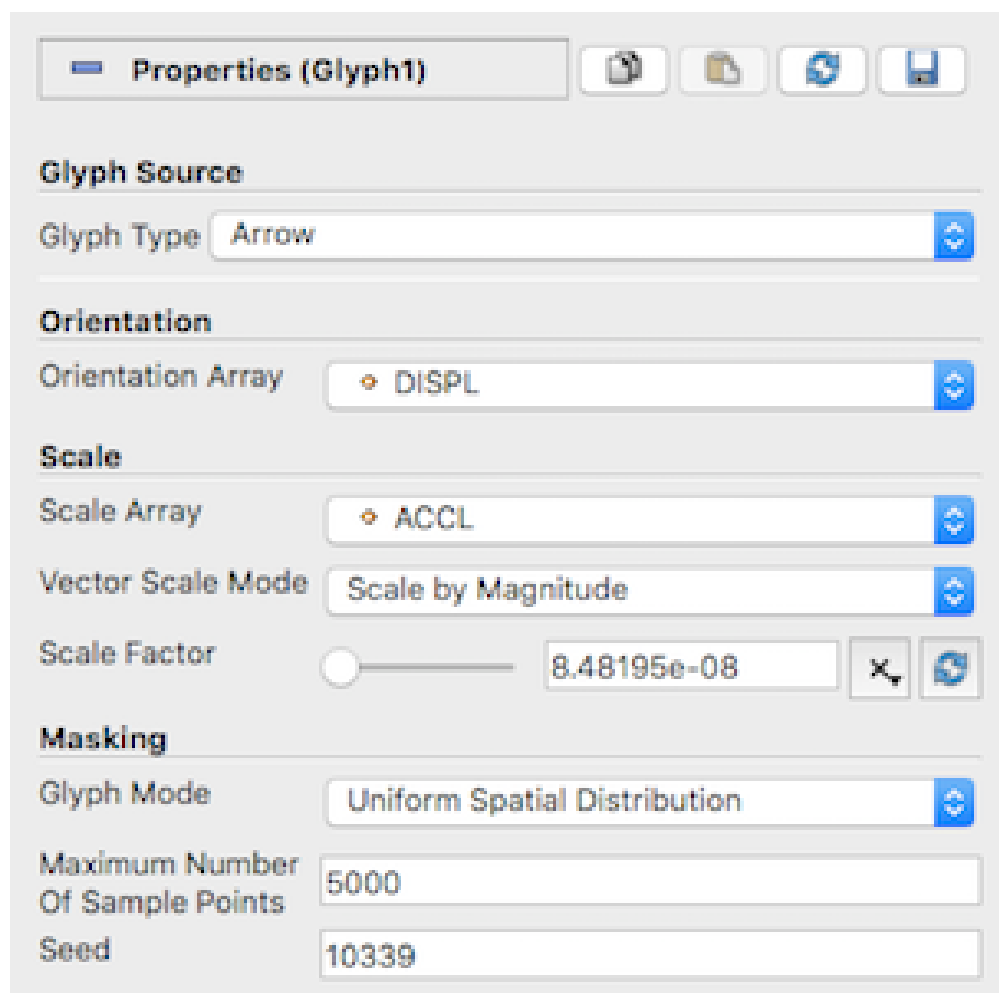


Fig. 1.66: The Properties panel for the Glyph filter.

The Masking properties control which points from the input dataset get glyphed. The Glyph Mode controls how points are selected to be glyphs (Fig. 1.67). The available options are as follows:

- **All Points** : This selects all points in the input dataset for glyphing. Use this mode with caution and only when the input dataset has relatively few points. Since all points in the input dataset are glyphed, this can not only cause visual clutter, but also clog up memory and take a long time to generate and render the glyphs.
- **Every Nth Points** : This elects every n^{th} point in the input dataset for glyphing, where n can be specified using Stride . Setting Stride to 1 will have the same effect as All Points .
- **Uniform Spatial Distribution** : This selects a random set of points. The algorithm works by first computing up to Maximum Number of Sample Points in the space defined by the bounding box of the input dataset. Then, points in the input dataset that are close to the point in this set of sample points are glyphed. The Seed is used to seed the random number generator used to generate the sample points. This ensures that the random sample points are reproducible and consistent.

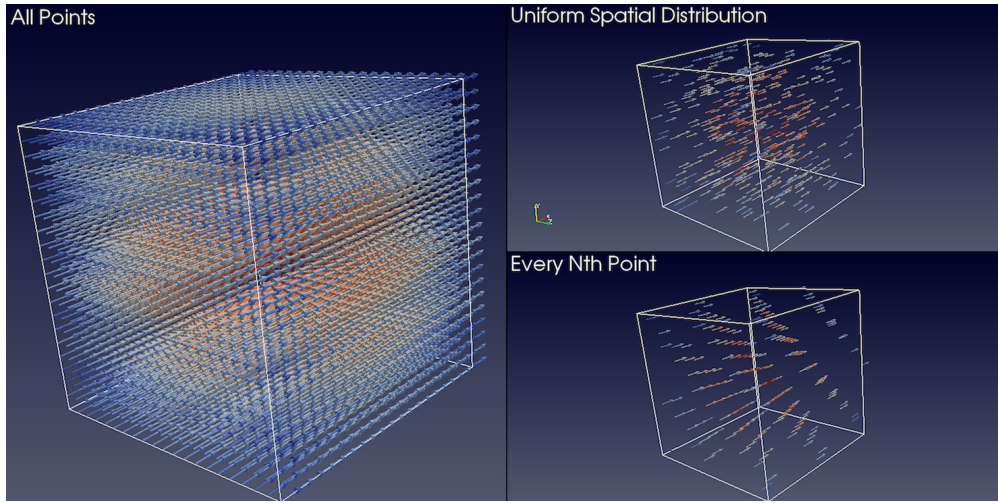


Fig. 1.67: Comparison between various Glyph Modes when applied to the same dataset generated by the Wavelet source.

Did you know?

The Glyph representation can be used for many of the same visualizations where a Glyph filter might be used. It may offer faster rendering and consume less memory than the Glyph filter with similar capabilities. In circumstances where generating a 3D geometry is required, e.g., when exporting glyph geometry to a file, the Glyph filter is required.

Glyph With Custom Source

Glyph With Custom Source is the same as Glyph , except that instead of a limited set of Glyph Type , you can select any data source producing a polygonal dataset (Section 1.3.1) available in the Pipeline Browser . To use this filter, select the data source you wish to glyph in the Pipeline Browser and attach this filter to it. You will be presented a dialog where you can set the Input (which defaults to the source you selected) and the Glyph Source .

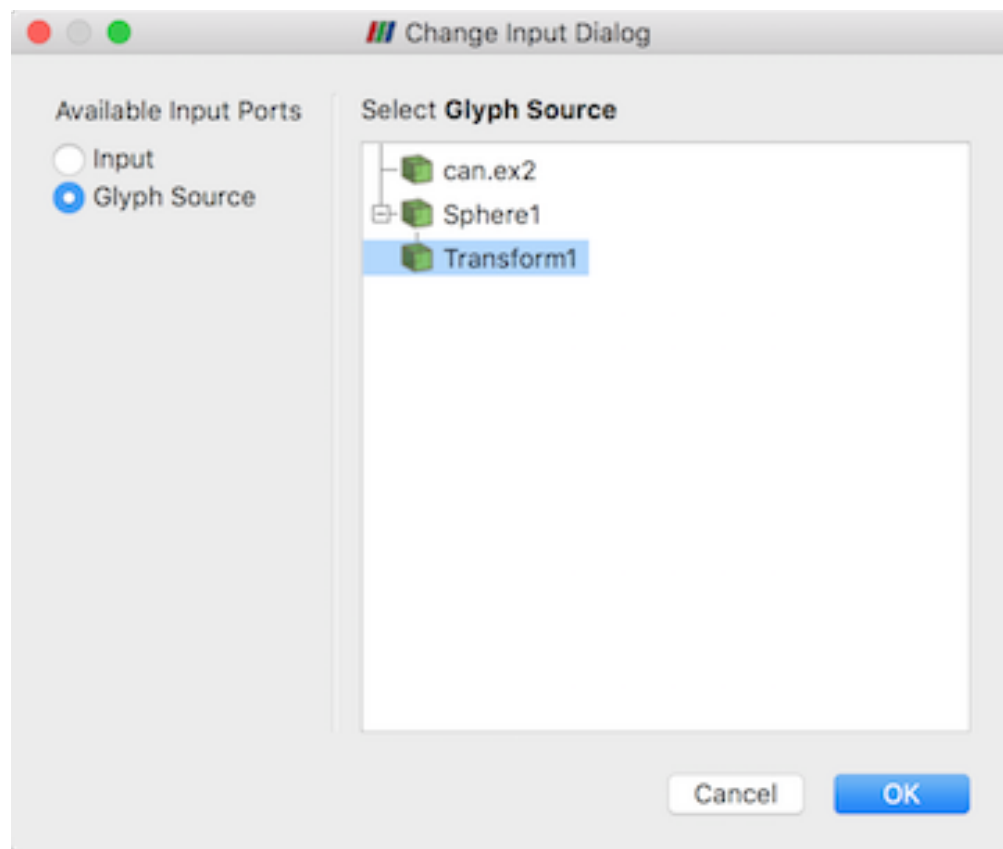


Fig. 1.68: Setting the Input and Glyph Source in the Glyph With Custom Source filter.

Stream Tracer

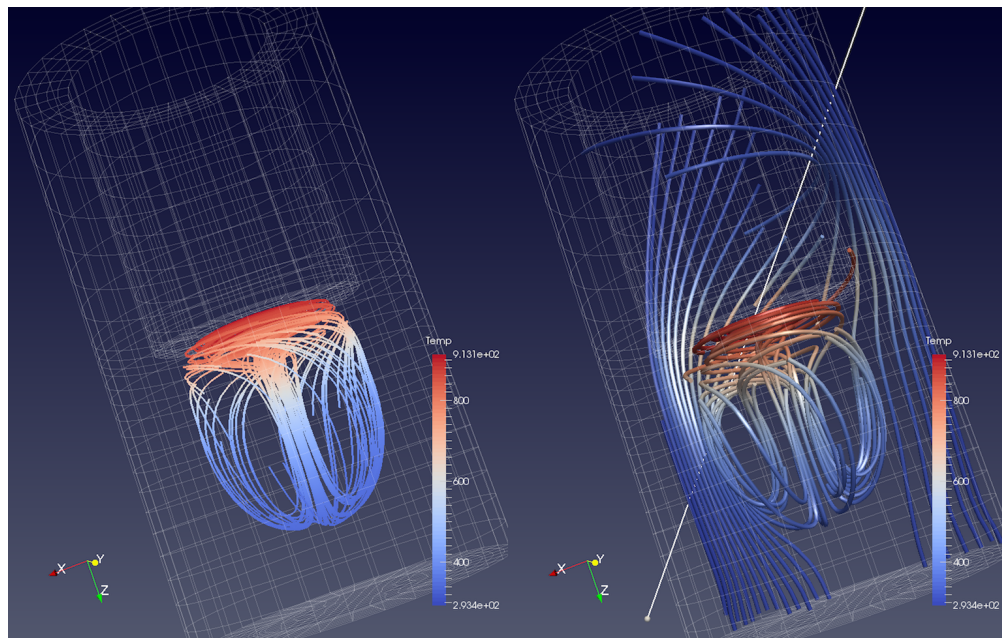


Fig. 1.69: Streamlines generated from the `disk_out_ref.ex2` dataset using the `Point Source` (left) and the `High Resolution Line Source` (right). On the left, we also added the `Tube` filter to the output of the `Stream Tracer` filter to generate 3D tubes rather than 1D polygonal lines, which can be hard to visualize due to lack of shading.

The `Stream Tracer` filter is used to generate streamlines for vector fields. In visualization, streamlines refer to curves that are instantaneously tangential to the the vector field in the dataset. They provide an indication of the direction in which the particles in the dataset would travel at that instant in time. The algorithm works by taking a set of points, known as *seed* points, in the dataset and then integrating the streamlines starting at these seed points.

In paraview, you can create this filter using the *Filters* menu, as well as the  button on the Common filters toolbar. To use this filter, you first select the attribute array to use as the `Vectors` for generating the streamline. `Integration Parameters` let you fine tune the streamline integration by specifying the direction to integrate, `Integration Direction`, as well as the type of integration algorithm to use, `Integrator Type`. Advanced integration parameters are available in the advanced view of the `Properties` panel that let you further tune the integration, including specifying the step size and others. You use the `Maximum Streamline Length` to limit the maximum length for the streamline – the longer the length, the longer the generated streamlines.

`Seeds` group lets you set how the seed points for generating the streamlines are produced. You have two options: `Point Source`, which produces a point cloud around the user-specified `Point` based on the parameters specified, and `High Resolution Line Source`, which produces seed points along the user-specified line. You can use the 3D widgets shown in the active `Render View` to interactively place the center for the point cloud or for defining the line.

Did you know?

The `Stream Tracer` filter produces a polydata with 1D lines for each of the generated streamlines. Since 1D lines cannot be shaded like surfaces in the `Render View`, you can get visualizations where it is hard to follow the streamlines. To give the streamlines some 3D structure, you can apply the `Tube` filter to the output of the streamlines. The properties on the `Tube` filter let you control the thickness of the tubes. You can also vary the thickness of the tubes based on data array, e.g., the magnitude of the vector field at the sample points in the streamline!

A script using the `Stream Tracer` filter in paraview typically looks like this:

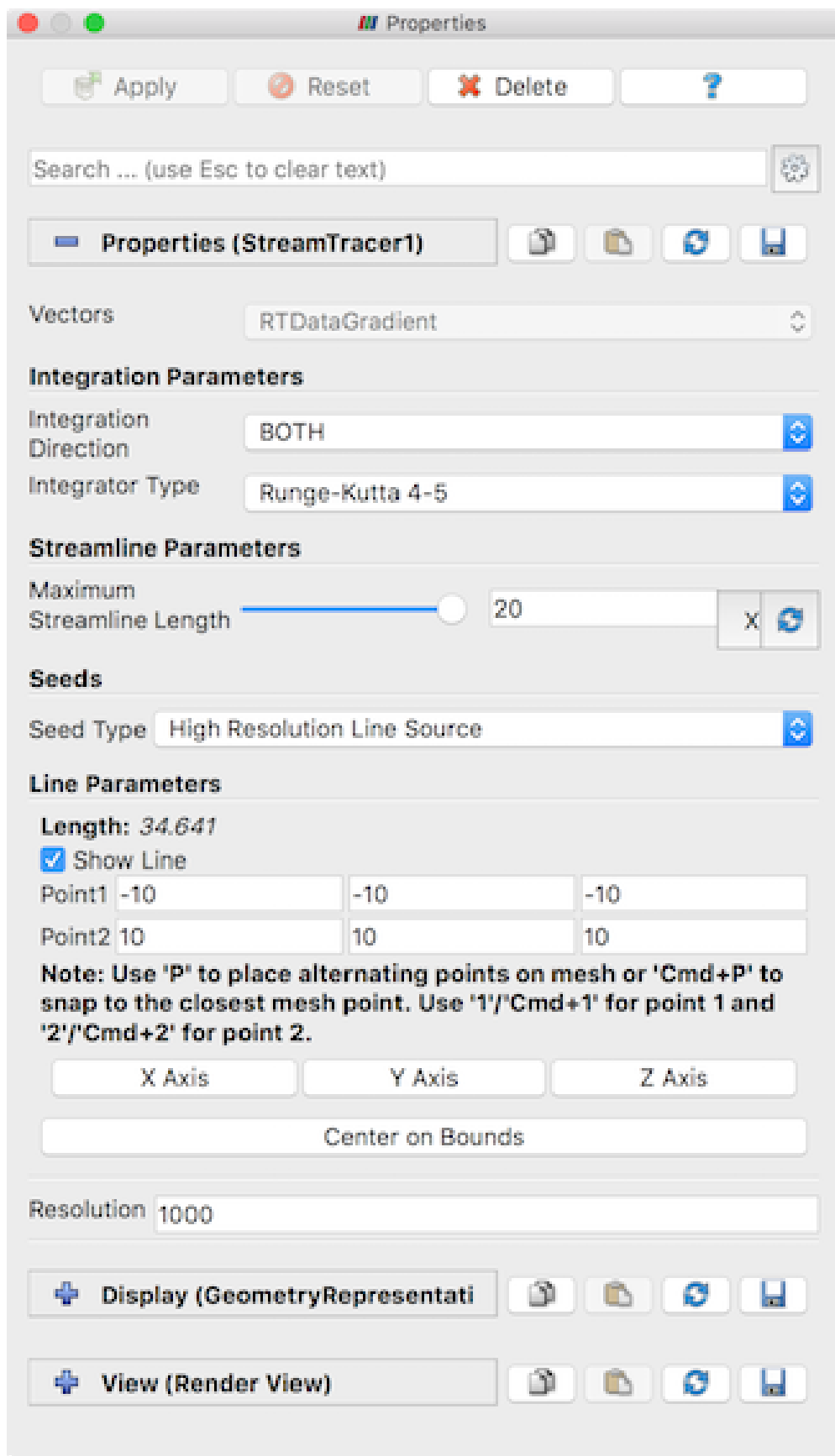


Fig. 1.70: The Properties panel showing the default properties for the Stream Tracer filter.

```
# find source
>>> disk_out_refex2 = FindSource('disk_out_ref.ex2')

# create a new 'Stream Tracer'
>>> streamTracer1 = StreamTracer(Input=disk_out_refex2,
                                  SeedType='Point Source')
>>> streamTracer1.Vectors = ['POINTS', 'V']

# init the 'Point Source' selected for 'SeedType'
>>> streamTracer1.SeedType.Center = [0.0, 0.0, 0.07999992370605469]
>>> streamTracer1.SeedType.Radius = 2.015999984741211

# show data in view
>>> Show()

# create a new 'Tube'
>>> tube1 = Tube(Input=streamTracer1)

# Properties modified on tube1
>>> tube1.Radius = 0.1611409378051758

# show the data from tubes in view
>>> Show()
```

Stream Tracer With Custom Source

Stream Tracer allows you to specify the seed points either as a point cloud or as a line source. However, if you want to provide your own seed points from another data producer, use the Stream Tracer With Custom Source . Similar to Glyph With Custom Source , this filter allows you to pick a second input connection to use as the seed points.

Resample With Dataset

Resample With Dataset samples the point and cell attributes of one dataset on to the points of another dataset. The two datasets are supplied to the filter using its two input ports: Input , which is the dataset that provides the attributes to resample, and Source , which is the dataset that provides the points to sample at. This filter is available under the *Filters* menu.

Resample To Image

Resample To Image is a specialization of Resample With Dataset . The filter takes one input and samples its point and cell attributes onto a uniform grid of points. The bounds and extents of the uniform grid can be specified using the properties panel. By default, the bounds are set to the bounds of the input dataset. The output of the filter is an Image dataset.

Some operations can be performed more efficiently on uniform grid datasets. Volume rendering is one such operation. The Resample to Image filter can be used to convert any dataset to Image data before performing such operations.

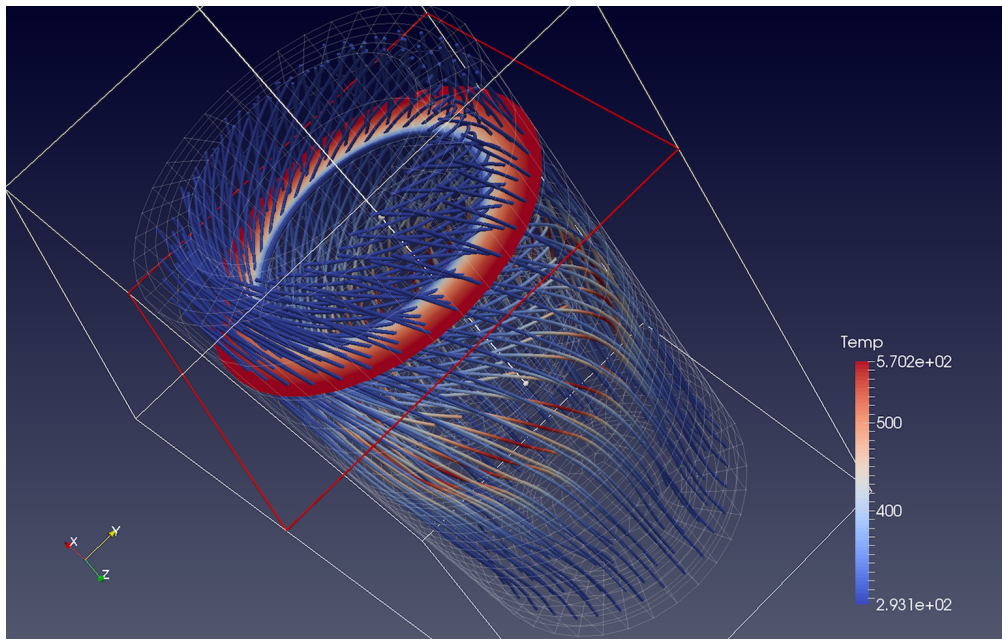


Fig. 1.71: Streamlines generated from the `disk_out_ref.ex2` dataset using the output of the `Slice` filter as the Source for seed points.

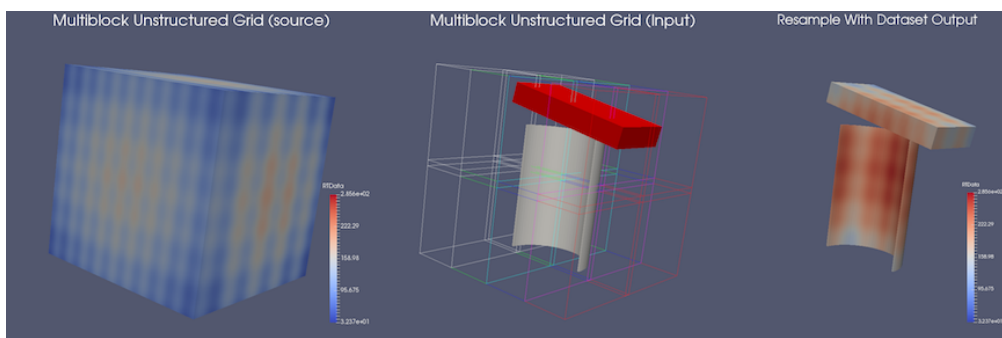


Fig. 1.72: An example of `Resample With Dataset`. On the left is a multiblock tetrahedra mesh (`Input`). The middle shows a multiblock unstructured grid (`Source`). The outline of `Input` is also shown in this view. The result of applying the filter is shown on the right

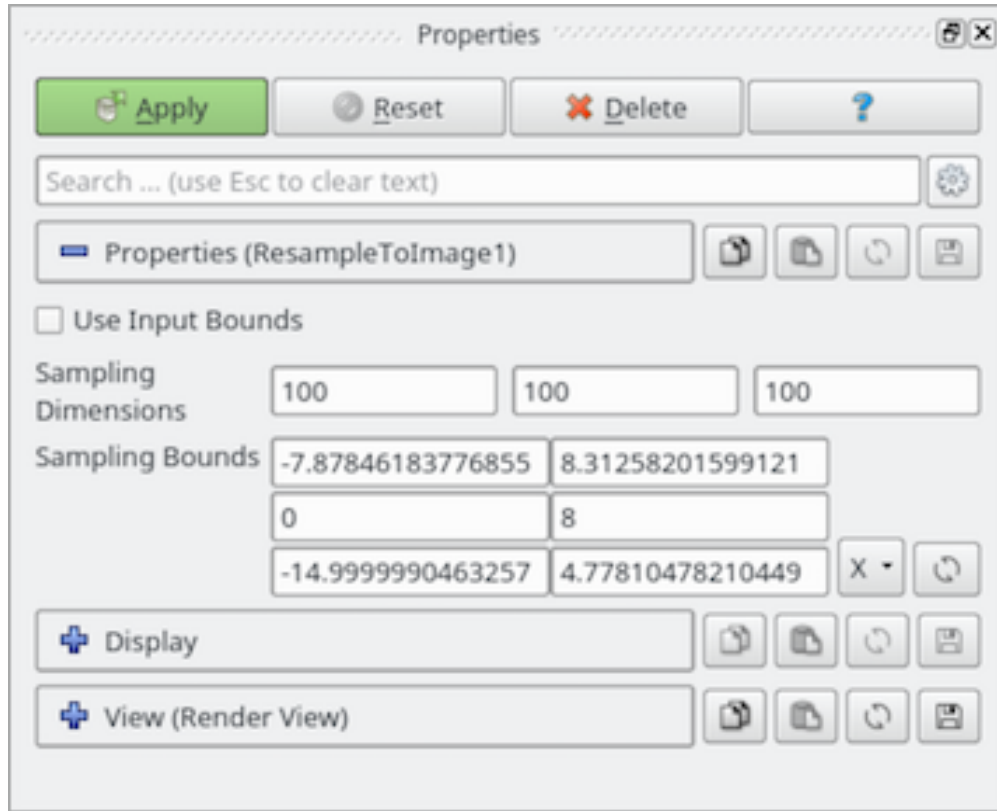


Fig. 1.73: The Properties panel for Resample To Image filter.

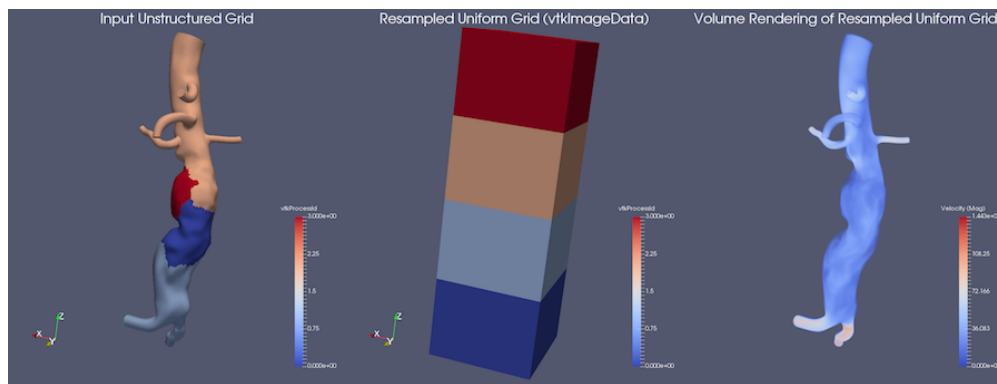


Fig. 1.74: An example of Resample To Image . The left portion shows the input (unstructured grid), and the middle displays the output image data. On the right is a volume rendering of the resampled data.

Probe

Probe samples the input dataset at a specific point location to obtain the cell data attributes for the cell containing the point as well as the interpolated point data attributes. You can either use the **SpreadSheet View** or the **Information** panel to inspect the probed values. The probe location can be specified using the interactive 3D widget shown in the active **Render View**.

Plot over line

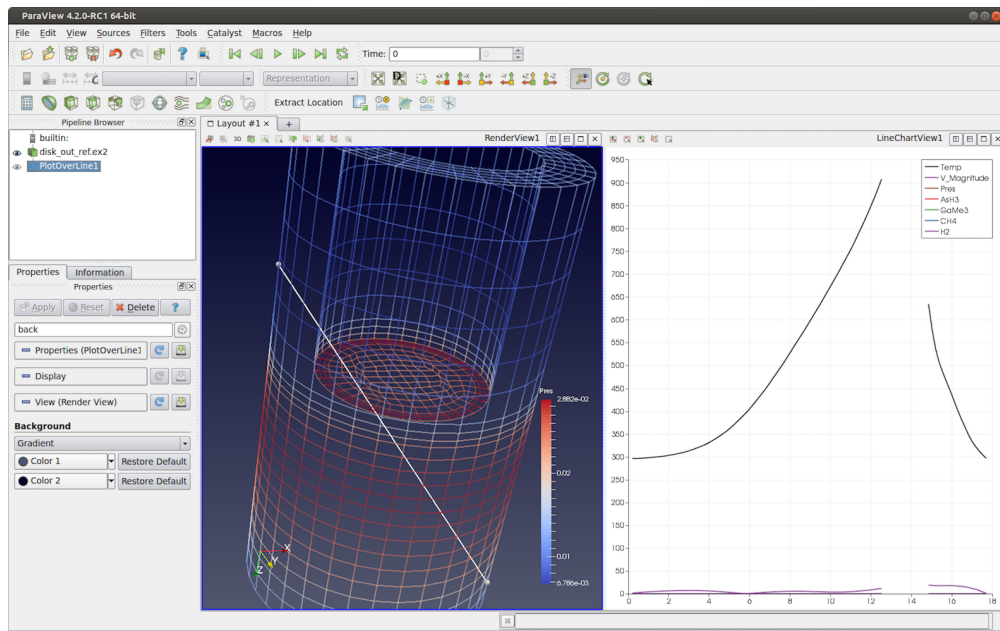


Fig. 1.75: The **Plot Over Line** filter applied to the `disk_out_ref.ex2` dataset to plot values at sampled locations along the line. Gaps in the line correspond to the locations in the input dataset where the line falls outside the dataset.

Plot Over Line will sample the input dataset along the specified line and then plot the results in **Line Chart View**. Internally, this filter uses the same mechanism as the **Probe** filter, probing along the points in the line to get the containing cell attributes and interpolated point attributes.

Using the **Resolution** property on the **Properties** panel, you can control the number of sample points along the line.

1.5.9 Filters for attribute manipulation

The filters covered in this section are used to add new attribute arrays to the dataset, which are typically used to add derived quantities to use in pipelines for further processing.

Calculator

The **Calculator** filter computes a new data array or new point coordinates as a function of existing input arrays. If point-centered arrays are used in the computation of a new data array, the resulting array will also be point-centered. Similarly, computations using cell-centered arrays will produce a new cell-centered array. If the function is computing point coordinates (requested by checking the **Coordinate Results** property on the **Properties** panel), the result of the function must be a three-component vector. The **Calculator** interface operates similarly to a scientific calculator. In creating the function to evaluate, the standard order of operations applies. Each of the calculator functions is described below. Unless otherwise noted, enclose the operand in parentheses using the (and) buttons.

- **Clear** : Erase the current function.
- **/**: Divide one scalar by another. The operands for this function are not required to be enclosed in parentheses.
- *****: Multiply two scalars, or multiply a vector by a scalar (scalar multiple). The operands for this function are not required to be enclosed in parentheses.
- **-**: Negate a scalar or vector (unary minus), or subtract one scalar or vector from another. The operands for this function are not required to be enclosed in parentheses.
- **+**: Add two scalars or two vectors. The operands for this function are not required to be enclosed in parentheses.
- **sin** : Compute the sine of a scalar.
- **cos** : Compute the cosine of a scalar.
- **tan** : Compute the tangent of a scalar.
- **asin** : Compute the arcsine of a scalar.
- **acos** : Compute the arccosine of a scalar.
- **atan** : Compute the arctangent of a scalar.
- **sinh** : Compute the hyperbolic sine of a scalar.
- **cosh** : Compute the hyperbolic cosine of a scalar.
- **tanh** : Compute the hyperbolic tangent of a scalar.
- **min** : Compute minimum of two scalars.
- **max** : Compute maximum of two scalars.
- **x^y** : Raise one scalar to the power of another scalar. The operands for this function are not required to be enclosed in parentheses.
- **sqrt** : Compute the square root of a scalar.
- **e^x** : Raise e to the power of a scalar.
- **log10** : Compute the logarithm of a scalar to the base 10.
- **ln** : Compute the logarithm of a scalar to the base e.
- **ceil** : Compute the ceiling of a scalar.
- **floor** : Compute the floor of a scalar.
- **abs** : Compute the absolute value of a scalar.

- `v1.v2` : Compute the dot product of two vectors. The operands for this function are not required to be enclosed in parentheses.
- `cross` : Compute cross product of two vectors.
- `mag` : Compute the magnitude of a vector.
- `norm` : Normalize a vector. The operands are described below. The digits 0-9 and the decimal point are used to enter constant scalar values.
- `iHat`, `jHat`, and `kHat` are vector constants representing unit vectors in the X, Y, and Z directions, respectively.

The Scalars menu lists the names of the scalar arrays and the components of the vector arrays of either the point-centered or cell-centered data. The Vectors menu lists the names of the point-centered or cell-centered vector arrays. The function will be computed for each point (or cell) using the scalar or vector value of the array at that point (or cell). The filter operates on any type of dataset, but the input dataset must have at least one scalar or vector array. The arrays can be either point-centered or cell-centered. The Calculator filter's output is of the same dataset type as the input.

A common use-case is to convert three input scalars into a vector array. For that, the Function would look something like: `scalarx * iHat + scalary * jHat + scalarz * kHat`.

The Properties panel provides access to several options for this filter. Checking Coordinate Results, Result Normals, or Result TCoords will set the computed array as the point coordinates, normals, or texture coordinates, respectively. Result Array Name is used to specify a name for the computed array. The default is Result.

Sometimes, the expression can yield invalid values. To replace all invalid values with a specific value, check the Replace Invalid Results checkbox and then enter the value to use to replace invalid values using the Replacement Value. The output array data type is set with the Result Array Type property.

Python calculator

The Python Calculator is similar to Calculator in that it processes one or more input arrays based on an expression provided by the user to produce a new output array. However, it uses Python (and NumPy) to do the computation. Therefore, it provides more expressive expression capabilities.

Specify the Expression to use, the Array Association to indicate the array association (Point Data or Cell Data), the name of output array (Array Name), and a toggle that controls whether the input arrays are copied to the output (Copy Array).

Basic tutorial

Start by creating a Sphere source and applying the Python Calculator to it. As the first expression, use the following and apply:

5

This should create an array name `result` in the output point data. Note that this is an array that has a value of 5 for each point. When the expression results in a single value, the calculator will automatically make a constant array. Next, try the following:

Normals

Now, the `result` array should be the same as the input array Normals. As described in detail later, various functions are available through the calculator. For example, the following is a valid expression:

`sin(Normals) + 5`

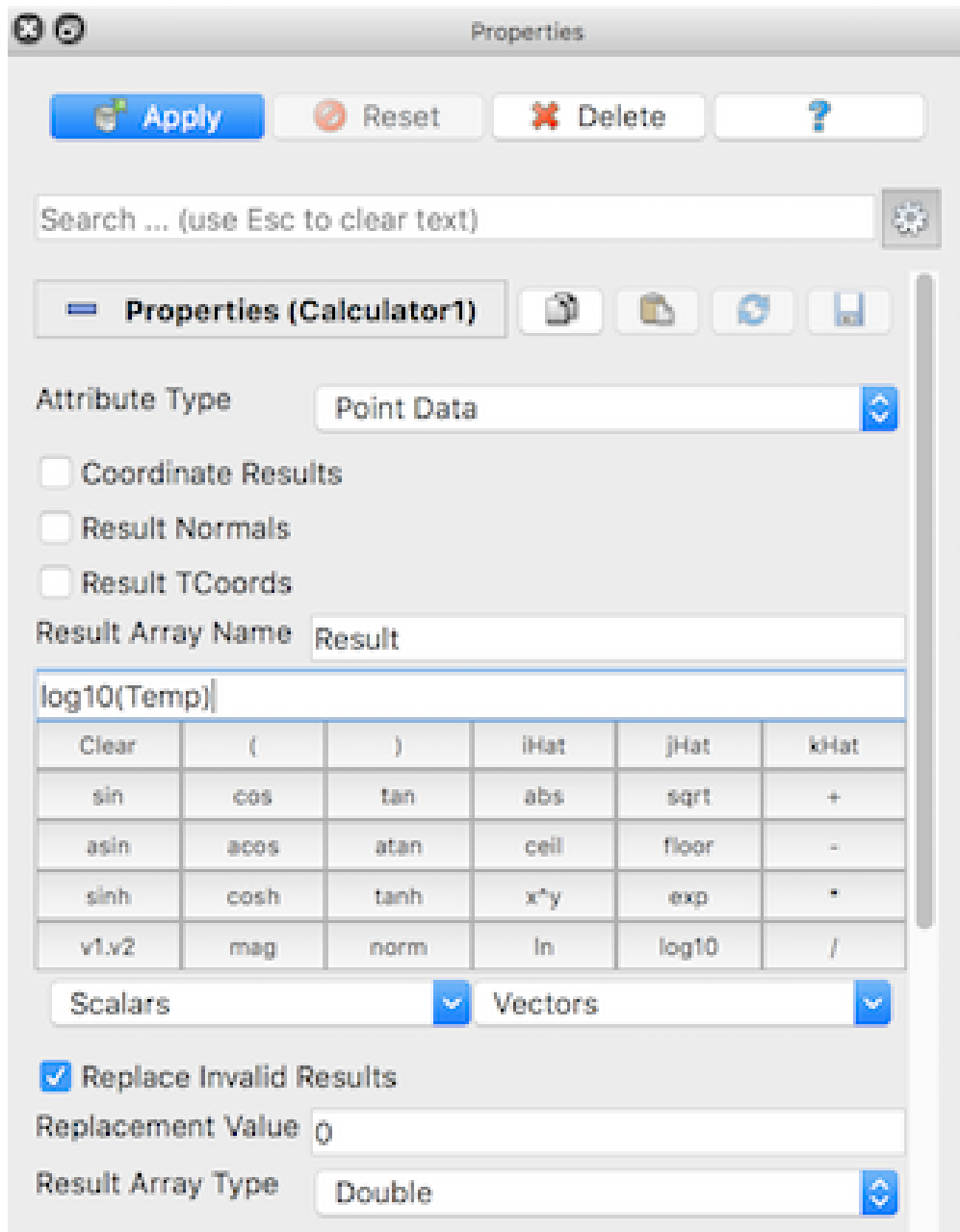


Fig. 1.76: The Properties panel for the Calculator filter showing the advanced properties.

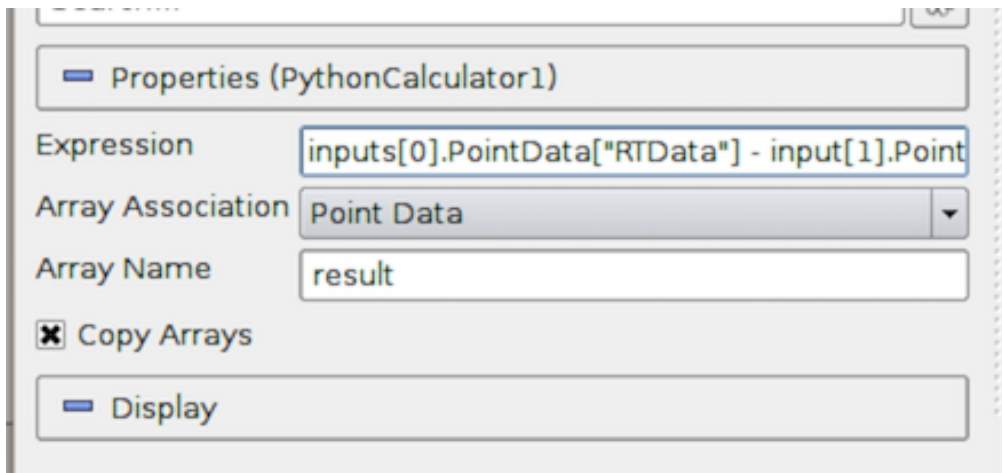


Fig. 1.77: The Properties Panel for Python Calculator.

It is very important to note that the Python Calculator has to produce one value per point or cell depending on the Array Association parameter. Most of the functions described here apply individually to all point or cell values and produce an array the same dimensions as the input. However, some of them, such as `min()` and `max()`, produce single values.

Common Errors

In the Programmable Filter, all the functions in `vtk.numpy_interface.algorithms` are imported prior to executing the script. As a result, some built-in functions, such as `min` and `max`, are clobbered by that import. To use the built-in functions, import the `__builtin__` module and access those functions with, e.g., `__builtin__.min` and `__builtin__.max`

Accessing data

There are several ways of accessing input arrays within expressions. The simplest way is to access it by name:

```
sin(Normals) + 5
```

This is equivalent to:

```
sin(inputs[0].PointData['Normals']) + 5
```

The example above requires some explanation. Here, `inputs[0]` refer to the first input (dataset) to the filter. Python Calculator can accept multiple inputs. Each input can be accessed as `inputs[0]`, `inputs[1]`, ... You can access the point or cell data of an input using the `.PointData` or `.CellData` qualifiers. You can then access individual arrays within the point or cell data containers using the `[]` operator. Make sure to use quotes or double-quotes around the array name. Arrays that have names with certain characters (such as space, +, -, *, /) can only be accessed using this method.

Certain functions apply directly on the input mesh. These filters expect an input dataset as argument. For example,

```
area(inputs[0])
```

For data types that explicitly define the point coordinates, you can access the coordinates array using the `.Points` qualifier. The following extracts the first component of the coordinates array:

```
inputs[0].Points[:,0]
```

Note that for certain data types, mainly image data (uniform rectilinear grids) and rectilinear grids, point coordinates are defined implicitly and cannot be accessed as an array.

Comparing multiple datasets

The Python Calculator can be used to compare multiple datasets, as shown by the following example.

- Go to the Menu Bar, and select *File > Disconnect* to clear the Pipeline.
- Select *Source > Mandelbrot*, and then click *Apply*, which will set up a default version of the Mandelbrot Set. The data for this set are stored in a 251×251 scalar array.
- Select *Source > Mandelbrot* again, and then go to the *Properties* panel and set the Maximum Number of Iterations to 50. Click *Apply*, which will set up a different version of the Mandelbrot Set, represented by the same size array.
- Hold the Shift key down and select both of the Mandelbrot entries in the Pipeline Inspector, and then go to the Menu Bar, and select *Filter > Python Calculator*. The two Mandelbrot entries will now be shown as linked, as inputs, to the Python Calculator .
- In the Properties panel for the Python Calculator filter, enter the following into the Expression box:

```
inputs[1].PointData['Iterations'] - inputs[0].PointData['Iterations']
```

This expression specifies the difference between the second and the first Mandelbrot arrays. The result is saved in a new array called **results** . The prefixes in the names for the array variables, **inputs[1]** and **inputs[0]** , refer to the first and second Mandelbrot entries, respectively, in the Pipeline. **PointData** specifies that the inputs contain point values. The quoted label '**Iterations**' is the local name for these arrays. Click *Apply* to initiate the calculation.

Click the **Display** tab in the **Properties Panel** for the Python Calculator , and go to the first tab to the right of the *Color by* label. Select the item **results** in that tab, which will cause the display window to the right to show the results of the expression we entered in the Python Calculator . The scalar values representing the difference between the two Mandelbrot arrays are represented by colors that are set by the current color map (see *Edit Color Map...* for details).

There are a few things to note:

- Python Calculator will always copy the mesh from the first input to its output.
- All operations are applied point-by-point. In most cases, this requires that the input meshes (topology and geometry) are the same. At the least, it requires that the inputs have the same number of points and cells.
- In parallel execution mode, the inputs have to be distributed exactly the same way across processes.

Basic Operations

The Python Calculator supports all of the basic arithmetic operations using the **+**, **-**, ***** and **/** operators. These are always applied element-by-element to point and cell data including scalars, vectors, and tensors. These operations also work with single values. For example, the following adds 5 to all components of all Normals.

```
Normals + 5
```

The following adds 1 to the first component, 2 to the second component, and 3 to the third component:

```
Normals + [1,2,3]
```

This is specially useful when mixing functions that return single values. For example, the following normalizes the Normals array:

```
(Normals - min(Normals))/(max(Normals) - min(Normals))
```

A common use case in a calculator is to work on one component of an array. This can be accomplished with the following:

```
Normals[:, 0]
```

The expression above extracts the first component of the Normals vector. Here, `:` is a placeholder for “all elements”. One element can be extracted by replacing `:` with an index. For example, the following creates a constant array from the first component of the normal of the first point:

```
Normals[0, 0]
```

Alternatively, the following assigns the normal of the first point to all points:

```
Normals[0, :]
```

It is also possible to create a vector array from two or three scalar arrays using the `make_vector()` function:

```
make_vector(velocity_x, velocity_y, velocity_z)
```

For temporal datasets, you also have access to the dataset timestep index or time value in the expression as `t_index` or `time_index`, and `t_value` or `time_value` respectively. When dealing with multiple inputs, you can specify the same variable names scoped on the appropriate input e.g. `inputs[0].t_index`.

The locations of points are available in the `Points` variable for datasets that define explicit points positions.

In some datasets, field data is used to store global data values not associated with cells or points. To use field data in a Python `Calculator` expression, access it with the `FieldData` dictionary available in the input as in the following example:

```
VolumeOfCell * inputs[0].FieldData['MaterialData'][time_index]
```

Did you know?

Under the cover, the Python `Calculator` uses NumPy. All arrays in the expression are compatible with NumPy arrays and can be used where NumPy arrays can be used. For more information on what you can do with these arrays, consult with the NumPy references:cite: numpy.

Functions

The following is a list of functions available in the `Python Calculator`. Note that this is a partial list, since most of the NumPy and SciPy functions can be used in the `Python Calculator`. Many of these functions can take single values or arrays as argument.

- `abs(x)` : Returns the absolute value(s) of x .
- `add(x, y)` : Returns the sum of two values. x and y can be single values or arrays. This is the same as $x + y$.
- `area(dataset)` : Returns the surface area of each cell in a mesh.
- `aspect(dataset)` : Returns the aspect ratio of each cell in a mesh.
- `aspect_gamma(dataset)` : Returns the aspect ratio gamma of each cell in a mesh.
- `condition(dataset)` : Returns the condition number of each cell in a mesh.
- `cross(x, y)` : Returns the cross product for two 3D vectors from two arrays of 3D vectors.
- `curl(array)` : Returns the curl of an array of 3D vectors.
- `divergence(array)` : Returns the divergence of an array of 3D vectors.
- `divide(x, y)` : Element-by-element division. x and y can be single values or arrays. This is the same as `math:frac{x}{y}`.
- `det(array)` : Returns the determinant of an array of 2D square matrices.
- `determinant(array)` : Returns the determinant of an array of 2D square matrices.
- `diagonal(dataset)` : Returns the diagonal length of each cell in a dataset.
- `dot(a1, a2)` : Returns the dot product of two scalars/vectors of two array of scalars/vectors.
- `eigenvalue(array)` : Returns the eigenvalue of an array of 2D square matrices.
- `eigenvector(array)` : Returns the eigenvector of an array of 2D square matrices.
- `exp(x)` : Returns e^x .
- `gradient(array)` : Returns the gradient of an array of scalars or vectors.
- `inv(array)` : Returns the inverse an array of 2D square matrices.
- `inverse(array)` : Returns the inverse of an array of 2D square matrices.
- `jacobian(dataset)` : Returns the jacobian of an array of 2D square matrices.
- `laplacian(array)` : Returns the jacobian of an array of scalars.
- `ln(array)` : Returns the natural logarithm of an array of scalars/vectors/tensors.
- `log(array)` : Returns the natural logarithm of an array of scalars/vectors/tensors.
- `log10(array)` : Returns the base 10 logarithm of an array of scalars/vectors/tensors.
- `make_point_mask_from_NaNs(dataset, array)` : This function will create a ghost array corresponding to an input with NaN values. For each NaN value, the output array will have a corresponding value of `vtk.vtkDataSetAttributes.HIDDENPOINT`. These values are also combined with any ghost values that the dataset may have.
- `make_cell_mask_from_NaNs(dataset, array)` : This function will create a ghost array corresponding to an input with NaN values. For each NaN value, the output array will have a corresponding value of `vtk.vtkDataSetAttributes.HIDENCELL`. These values are also combined with any ghost values that the dataset may have.

- **max(array)** : Returns the maximum value of the array as a single value. In parallel, compute the max accross processes.
- **max_angle(dataset)** : Returns the maximum angle of each cell in a dataset.
- **mag(a)** : Returns the magnitude of an array of scalars/vectors.
- **mean(array)** : Returns the mean value of an array of scalars/vectors/tensors. In parallel, compute the mean accross processes.
- **min(array)** : Returns the minimum value of the array as a single value. In parallel, compute the min accross processes.
- **min_angle(dataset)** : Returns the minimum angle of each cell in a dataset.
- **mod(x, y)** : Same as remainder (x, y).
- **multiply(x, y)** : Returns the product of x and y . x and y can be single values or arrays. Note that this is an element-by-element operation when x and y are both arrays. This is the same as $x \times y$.
- **negative(x)** : Same as $-x$.
- **norm(a)** : Returns the normalized values of an array of scalars/vectors.
- **power(x, a)** : Exponentiation of x with a . Here, both x and a can either be a single value or an array. If x and y are both arrays, a one-by-one mapping is used between two arrays.
- **reciprocal(x)** : Returns $\frac{1}{x}$.
- **remainder(x, y)** : Returns $x - y \times \text{floor}(\frac{x}{y})$. x and y can be single values or arrays.
- **rint(x)** : Rounds x to the nearest integer(s).
- **shear(dataset)** : Returns the shear of each cell in a dataset.
- **skew(dataset)** : Returns the skew of each cell in a dataset.
- **square(x)** : Returns $x * x$.
- **sqrt(x)** : Returns $\sqrt[2]{x}$.
- **strain(array)** : Returns the strain of an array of 3D vectors.
- **subtract(x, y)** : Returns the difference between two values. x and y can be single values or arrays. This is the same as $x - y$.
- **surface_normal(dataset)** : Returns the surface normal of each cell in a dataset.
- **trace(array)** : Returns the trace of an array of 2D square matrices.
- **volume(dataset)** : Returns the volume normal of each cell in a dataset.
- **vorticity(array)** : Returns the vorticity/curl of an array of 3D vectors.
- **vertex_normal(dataset)** : Returns the vertex normal of each point in a dataset.

Trigonometric Functions

Below is a list of supported trigonometric functions:

$\sin(x)$	$\arccos(x)$	$\cosh(x)$	$\operatorname{arctanh}(x)$
$\cos(x)$	$\arctan(x)$	$\tanh(x)$	
$\tan(x)$	$\operatorname{hypot}(x_1, x_2)$	$\operatorname{arcsinh}(x)$	
$\arcsin(x)$	$\sinh(x)$	$\operatorname{arccosh}(x)$	

Gradient

The Gradient filter computes the gradient of a cell or point data array for any type of dataset.

For unstructured grids, the gradient for cell data corresponds to the cell derivatives. For point data, the gradient at a given point is computed as the average of the derivatives of the cells to which the point belongs.

For structured grids, the gradient is computed using central differencing, except on the boundary of the dataset where forward and backward differencing is used for the boundary elements.

This filter can optionally compute the divergence, vorticity (also known as the curl), and Q-criterion. A 3-component array is required in order to compute these quantities. By default, only the gradient computation is enabled.

In the case of a uniform rectilinear grid (see [Section 1.3.1](#)), a specific implementation which efficiently computes the gradient of point data arrays is also available. This implementation extends the use of central differencing on the boundary elements after duplication of the boundary values. To activate this option, set the Boundary Method property to Smoothed, as shown in [Fig. 1.78](#).

Mesh Quality

The Mesh Quality filter creates a new cell array containing a geometric measure of each cell's fitness. Different quality measures can be chosen for different cell shapes.

Triangle Quality indicates which quality measure will be used to evaluate triangle quality. The Radius Ratio is the size of a circle circumscribed by a triangle's three vertices divided by the size of a circle tangent to a triangle's three edges. The Edge Ratio is the ratio of the longest edge length to the shortest edge length.

Quad Quality indicates which quality measure will be used to evaluate quad cells.

Tet Quality indicates which quality measure will be used to evaluate tetrahedral quality. The Radius Ratio is the size of a sphere circumscribed by a tetrahedron's four vertices divided by the size of a circle tangent to a tetrahedron's four faces. The Edge Ratio is the ratio of the longest edge length to the shortest edge length. The Collapse Ratio is the minimum ratio of height of a vertex above the triangle opposite it, divided by the longest edge of the opposing triangle across all vertex/triangle pairs.

HexQualityMeasure indicates which quality measure will be used to evaluate quality of hexahedral cells.

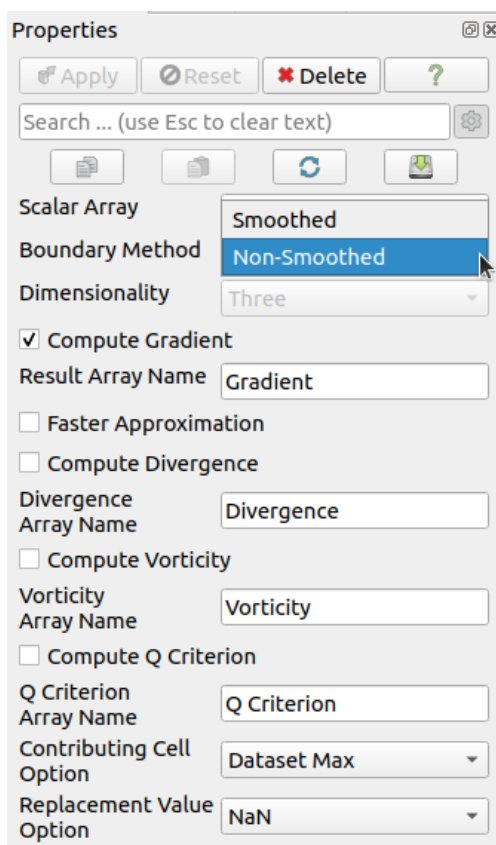


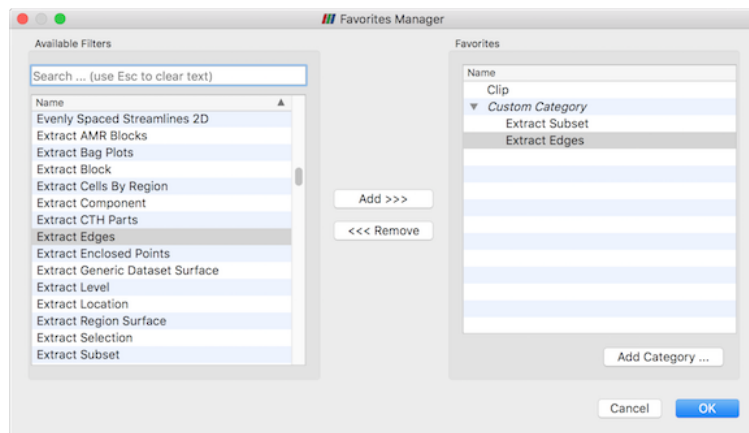
Fig. 1.78: The Properties Panel for the Gradient filter applied to a uniform structured grid.

1.5.10 White-box filters

This includes the **Programmable Filter** and **Programmable Source**. For these filters/sources, you can add Python code to do the data generation or processing. We'll cover writing Python code for these in [Section 2.4](#).

1.5.11 Favorite filters

If you use some filters more than others, you can organize them in the *Filters > Favorites* menu. This can be done from the context menu in the pipeline or through the *Filters > Manage Favorites* menu as shown in [Fig. 1.79](#). In this dialog you can create categories and subcategories. It supports drag'n'drop operation to sort and move filters and categories. Moreover, **Favorites** are highlighted in the other filter submenus on supported platforms. Favorites are saved in user settings so they can be used in other subsequent ParaView sessions.



[Fig. 1.79:](#) The Favorites Manager dialog. Left: the list of available filters. Right: the favorites, organized into categories.

1.5.12 Best practices

Avoiding data explosion

The pipeline model that **ParaView** presents is very convenient for exploratory visualization. The loose coupling between components provides a very flexible framework for building unique visualizations, and the pipeline structure allows you to tweak parameters quickly and easily.

The downside of this coupling is that it can have a larger memory footprint. Each stage of this pipeline maintains its own copy of the data. Whenever possible, **ParaView** performs shallow copies of the data so that different stages of the pipeline point to the same block of data in memory. However, any filter that creates new data or changes the values or topology of the data must allocate new memory for the result. If **ParaView** is filtering a very large mesh, inappropriate use of filters can quickly deplete all available memory. Therefore, when visualizing large datasets, it is important to understand the memory requirements of filters.

Please keep in mind that the following advice is intended only for when dealing with very large amounts of data and the remaining available memory is low. When you are not in danger of running out of memory, the following advice is not relevant.

When dealing with structured data, it is absolutely important to know what filters will change the data to unstructured. Unstructured data has a much higher memory footprint, per cell, than structured data because the topology must be explicitly written out. There are many filters in **ParaView** that will change the topology in some way, and these filters will write out the data as an unstructured grid, because that is the only dataset that will handle any type of topology

that is generated. The following list of filters will write out a new unstructured topology in its output that is roughly equivalent to the input. These filters should never be used with structured data and should be used with caution on unstructured data.

<i>Append Datasets</i>	<i>Extract Edges</i>	<i>Subdivide</i>
<i>Append Geometry</i>	<i>Linear Extrusion</i>	<i>Tessellate</i>
<i>Clean</i>	<i>Loop Subdivision</i>	<i>Tetrahedralize</i>
<i>Clean to Grid</i>	<i>Reflect</i>	<i>Triangle Strips</i>
<i>Connectivity</i>	<i>Rotational Extrusion</i>	<i>Triangulate</i>
<i>D3</i>	<i>Shrink</i>	
<i>Delaunay 2D/3D</i>	<i>Smooth</i>	

Technically, the **Ribbon** and **Tube** filters should fall into this list. However, as they only work on 1D cells in poly data, the input data is usually small and of little concern.

This similar set of filters also outputs unstructured grids, but also tends to reduce some of this data. Be aware though that this data reduction is often smaller than the overhead of converting to unstructured data. Also note that the reduction is often not well balanced. It is possible (often likely) that a single process may not lose any cells. Thus, these filters should be used with caution on unstructured data and extreme caution on structured data.

<i>Clip</i>	<i>Extract Selection</i>
<i>Decimate</i>	<i>Quadric Clustering</i>
<i>Extract Cells by Region</i>	<i>Threshold</i>

Similar to the items in the preceding list, **Extract Subset** performs data reduction on a structured dataset, but also outputs a structured dataset. So the warning about creating new data still applies, but you do not have to worry about converting to an unstructured grid.

This next set of filters also outputs unstructured data, but it also performs a reduction on the dimension of the data (for example 3D to 2D), which results in a much smaller output. Thus, these filters are usually safe to use with unstructured data and require only mild caution with structured data.

<i>Cell Centers</i>	<i>Feature Edges</i>
<i>Contour</i>	<i>Mask Points</i>
<i>Extract CTH Fragments</i>	<i>Outline (curvilinear)</i>
<i>Extract CTH Parts</i>	<i>Slice</i>
<i>Extract Surface</i>	<i>Stream Tracer</i>

The filters below do not change the connectivity of the data at all. Instead, they only add field arrays to the data. All the existing data is shallow copied. These filters are usually safe to use on all data.

<i>Block Scalars</i>	<i>Octree Depth Scalars</i>
<i>Calculator</i>	<i>Point Data to Cell Data</i>
<i>Cell Data to Point Data</i>	<i>Process Id Scalars</i>
<i>Curvature</i>	<i>Random Vectors</i>
<i>Elevation</i>	<i>Resample with dataset</i>
<i>Generate Surface Normals</i>	<i>Surface Flow</i>
<i>Gradient</i>	<i>Surface Vectors</i>
<i>Level Scalars</i>	<i>Texture Map to...</i>
<i>Median</i>	<i>Transform</i>
<i>Mesh Quality</i>	<i>Warp (scalar)</i>
<i>Octree Depth Limit</i>	<i>Warp (vector)</i>

This final set of filters either add no data to the output (all data of consequence is shallow copied) or the data they add is generally independent of the size of the input. These are almost always safe to add under any circumstances (although they may take a lot of time).

<i>Annotate Time</i>	<i>Outline</i>
<i>Append Attributes</i>	<i>Outline Corners</i>
<i>Extract Block</i>	<i>Plot Global Variables Over Time</i>
<i>Extract Datasets</i>	<i>Plot Over Line</i>
<i>Extract Level</i>	<i>Plot Selection Over Time</i>
<i>Glyph</i>	<i>Probe Location</i>
<i>Group Datasets</i>	<i>Temporal Shift Scale</i>
<i>Histogram</i>	<i>Temporal Snap-to-Time-Steps</i>
<i>Integrate Variables</i>	<i>Temporal Statistics</i>
<i>Normal Glyphs</i>	

There are a few special case filters that do not fit well into any of the previous classes. Some of the filters, currently `Temporal Interpolator` and `Particle Tracer`, perform calculations based on how data changes over time. Thus, these filters may need to load data for two or more instances of time, which can double or more the amount of data needed in memory. The `Temporal Cache` filter will also hold data for multiple instances of time. Keep in mind that some of the temporal filters such as the `Temporal Statistics` and the filters that plot over time may need to iteratively load all data from disk. Thus, it may take an impractically long amount of time even if does not require any extra memory.

The `Programmable Filter` is also a special case that is impossible to classify. Since this filter does whatever it is programmed to do, it can fall into any one of these categories.

Culling data

When dealing with large data, it is best to cull out data whenever possible and do so as early as possible. Most large data starts as 3D geometry and the desired geometry is often a surface. As surfaces usually have a much smaller memory footprint than the volumes that they are derived from, it is best to convert to a surface early on. Once you do that, you can apply other filters in relative safety.

A very common visualization operation is to extract isosurfaces from a volume using the `Contour` filter. The `Contour` filter usually outputs geometry much smaller than its input. Thus, the `Contour` filter should be applied early if it is to be used at all. Be careful when setting up the parameters to the `Contour` filter because it still is possible for it to generate a lot of data which can happen if you specify many isosurface values. High frequencies such as noise around an isosurface value can also cause a large, irregular surface to form.

Another way to peer inside of a volume is to perform a `Slice` on it. The `Slice` filter will intersect a volume with a plane and allow you to see the data in the volume where the plane intersects. If you know the relative location of an interesting feature in your large dataset, slicing is a good way to view it.

If you have little *a priori* knowledge of your data and would like to explore the data without the long memory and processing time for the full dataset, you can use the `Extract Subset` filter to subsample the data. The subsampled data can be dramatically smaller than the original data and should still be well load balanced. Of course, be aware that you may miss small features if the subsampling steps over them and that once you find a feature you should go back and visualize it with the full dataset.

There are also several features that can pull out a subset of a volume: `Clip`, `Threshold`, `Extract Selection`, and `Extract Subset` can all extract cells based on some criterion. Be aware, however, that the extracted cells are almost never well balanced; expect some processes to have no cells removed. All of these filters, with the exception of `Extract Subset`, will convert structured data types to unstructured grids. Therefore, they should not be used unless the extracted cells are of at least an order of magnitude less than the source data.

When possible, replace the use of a filter that extracts 3D data with one that will extract 2D surfaces. For example, if you are interested in a plane through the data, use the `Slice` filter rather than the `Clip` filter. If you are interested

in knowing the location of a region of cells containing a particular range of values, consider using the Contour filter to generate surfaces at the ends of the range rather than extract all of the cells with the Threshold filter. Be aware that substituting filters can have an effect on downstream filters. For example, running the Histogram filter after Threshold will have an entirely different effect than running it after the roughly equivalent Contour filter.

1.6 Selecting Data

A typical visualization process has two components: setting up the visualization scene and performing the analysis of the results to gain insight. It is not uncommon for this process to be iterative. Often, what you are looking for drives from what filters you should use to extract the relevant information from the input datasets and what views will best represent that data. One of the ways of evaluating the results is inspecting the data or probing into it by identifying elements of interest. **ParaView** data selection mechanisms are designed specifically for such use-cases. In this chapter, we take a closer look at various ways of selecting data in **ParaView** and making use of these selections for data analysis.

1.6.1 Understanding selection

Broadly speaking, selection refers to selecting elements (either cells, points, rows (in case of tabular datasets), etc.) from datasets. Since data is ingested into **ParaView** using readers or sources and transformed using filters, when you create a selection, you are selecting elements from the dataset produced as the output of source, filter, or any such pipeline module.

There are many ways to create selections. Several views provide means to create specific selections. For example, in the **SpreadSheet View**, which shows the data attributes as a spreadsheet, you can simply click on any row to *select* that row. You can, of course, use the **Shift** and **CTRL** (or **ALT**) keys to select multiple rows, as in typical spreadsheet-based applications.

While this seems like an exercise in futility, you are hardly achieving anything by highlighting rows in a spreadsheet. What transforms this into a key tool is the fact that selections are *linked* among views (whenever possible). Linked selection means that you select an element from a dataset in a specific view. All other views that are showing the *same* dataset will also highlight the selected elements.

To make this easier, let's try a quick demo:

Starting with a fresh **paraview** session, create a sample dataset using the **Sources > Wavelet** menu, and then click the **Apply** button. If you are using **paraview** with a default setup, that should result in a dataset outline being shown in the default **Render View**. Next, let's split the view and create **SpreadSheet View**. The **SpreadSheet View** will automatically show the data produced by the **Wavelet** source. Upon closer inspection of the header in the **SpreadSheet View**, we see that the view is showing the **Point Data** or point attributes associated with the dataset. Now we have the same dataset, the data produced by the **Wavelet** source, shown in two views. Now, highlight a few rows in the **SpreadSheet View** by clicking on them. As soon as you start selecting rows, the **Render View** will start highlighting some points in space as tiny magenta specks ([Fig. 1.80](#)). That's linked selection in action! What is happening is that, as you highlight rows in the **SpreadSheet View**, you are creating a selection for selecting points (since the view is showing **Point Data**) corresponding to the rows. Due to the linking of selections between views, any other view that is showing the dataset (in this case, the **Render View**) will also highlight the selected points.

Of course, if you want to select cells instead of points, switch the **SpreadSheet View** to show cells by flipping the **Attribute** combo-box to **Cell Data** and then highlight rows. The **Render View** will show the selected cells as a wireframe, rather than points.

Conversely, you could have created the selection in the **Render View**, and the **SpreadSheet View** will also highlight the selected elements. We will see how to create such selection later in this chapter.

The first thing to note is that, when you create a new selection, the existing selection is cleared. Thus, there is at most one active selection in the application at any given time. As we shall see, certain views provide ways of expanding on the existing selection.

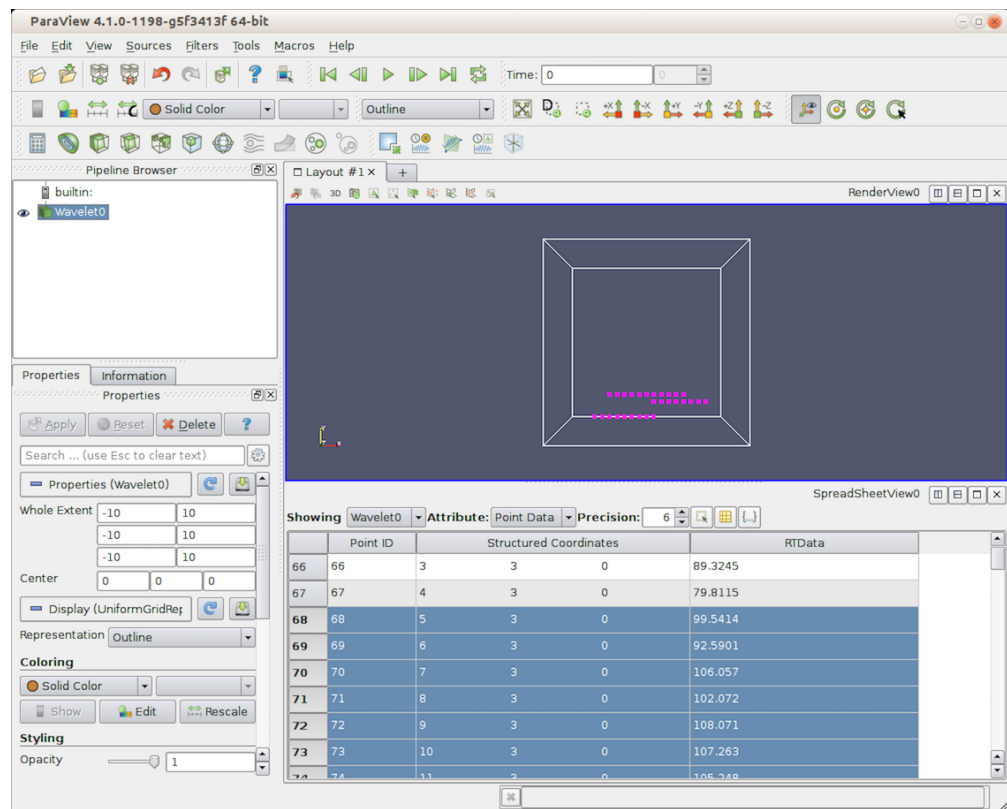


Fig. 1.80: Linked selection between views allows you to select elements in a view and view them in all other views showing the selected data. In this demo, as we select rows in the SpreadSheet View, the corresponding points in the 3D View get highlighted.

The second thing to note is that selections are *transient*, i.e., they cannot be undone/redone or saved in state files and loaded back. Nor can you apply filters or other transformation to the selections themselves. There are cases, however, where you may want to subset your dataset using the selection defined interactively and then apply filters and other analysis to that extracted subset. For that, there are filters available, namely **Extract Selection** and **Plot Selection Over Time**, that can capture the active selection in as the filter parameters and then produce a new dataset that is comprised of the selected elements.

The third thing to note is that there are different types of selections, e.g., id-based selections, where the selected elements are identified by their indices; frustum-based selections, where the selected elements are those that intersect a frustum defined in 3D space; query-based selections, where the selected elements are those that match the specified query string; and so on.

Did you know?

While, technically, selection is possible using a Python-based API, that API is currently cumbersome and requires understanding of some of **ParaView** internals. There are plans to simplify that API in future releases. As a result, we do not cover Python API for selection in this chapter. Once this new API is available, it will be included in this chapter.

1.6.2 Creating selections using views

Views provide a convenient mechanism for creating selections interactively. Views like **Render View** can create multiple types of selection (id- or frustum-based selections for selecting points and cells), while others like the **SpreadSheet View** and **Line Chart View** only support one type (id-based selections for points or cells).

Selecting in Render View

To create a selection in the **Render View**, you use the toolbar at the top of the view frame. There are two ways of selecting cells, points or blocks in **ParaView**: interactive and non-interactive.

ParaView enters a *non-interactive selection mode* when the user clicks one of the non-interactive selection buttons: The type of selection you are creating will depend on the button you clicked. Once in non-interactive selection mode, the cursor will switch to cross-hair and you can click and drag to create a selection region. Once you release the mouse, **ParaView** will attempt to create a selection for any elements in the selection region and will go back to default interaction mode.

To create a selection for cells visible in the view, use the button. For selecting visible points, use the button instead. Visible cells (and points) are only those cells (or points) that are currently rendered on the screen. Thus, elements that are occluded or are too small to be rendered on the screen will not be selected. If you want to select all data elements that intersect the view frustum formed by the selection rectangle you drew on the screen, use the button (or for points). In this case, all elements, visible or otherwise, that are within the 3D space defined by the selection frustum are selected.

To create a selection for blocks visible in the view use the button. Note that there is no frustum selection for blocks.

While most selection modes allow you to define the selection region as a rectangle, (and for points) enables you to define the selection region as a closed polygon. However, this is limited to surface elements (i.e., no frustum-based selection).

ParaView enters an *interactive selection mode* when the user clicks on one of the interactive selection buttons: In interactive selection mode, the user acts on visible elements (cells or points). **ParaView** highlights elements of the dataset as a user moves the cursor over them. An element can be selected by clicking on it. Clicking repeatedly on different elements adds them to the selection. A user ends the interactive selection mode by clicking on the interactive

selection button pushed in or by pressing the Esc key. This mode is also ended when the user enters a non-interactive selection mode. You can use button to interactively select cells of the dataset and use button to interactively select points.

When there are selected elements, the button can be used to clear the selection.

Several of these buttons have hotkeys too, such as S for visible cell selection, D for visible points selection, F for frustum-based cell selection, and G for frustum-based point selection. If you notice, these keys are right next to each other on the keyboard, starting with S, and are in the same order as the toolbar buttons themselves.

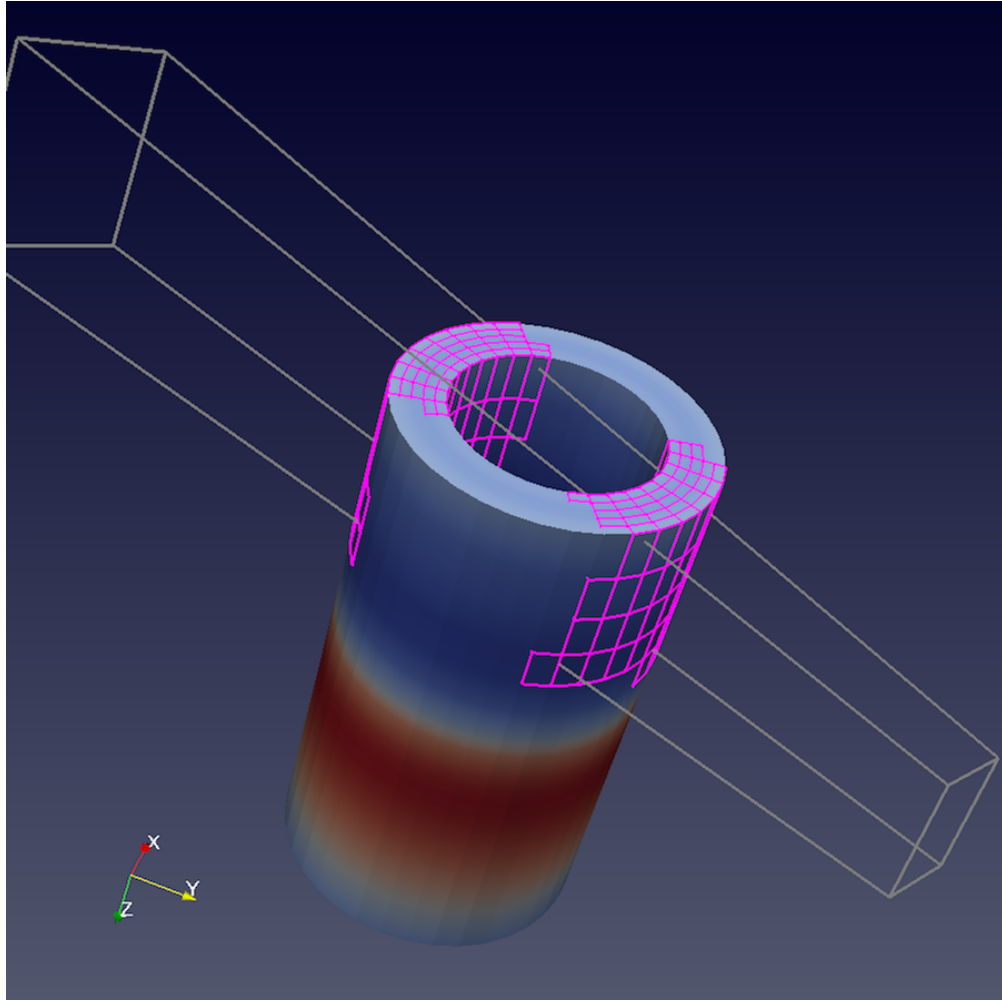


Fig. 1.81: Result of a frustum cell selection on disk_out_ref.ex2 dataset showing the frustum used to identify selected cells. All cells that fall in that frustum or that intersect it are selected, irrespective of whether they were visible from the view angle when the selection was made.

Did you know?

For id-based visible cell (or point) selection, you can expand the current selection by keeping the CTRL (or) key pressed when clicking and dragging in selection mode. paraview will then try to add to the current selection by adding the newly selected elements to the existing one. You can also subtract from the current selection using the , or even toggle using CTRL (or) + . Selection modifier buttons, in the toolbar, can be used for the same effect. Add : , remove :



, toggle : . These modifiers do not work, however, if the selection type (or the selected data) is different from the current selection. If so, the current selection will be cleared (as is the norm) and then the new selection will be created.

Selecting in SpreadSheet View

To create a selection in the `SpreadSheet View`, you simply click on the corresponding rows in the spreadsheet. You can use the `CTRL` (or `Shift`) keys to add to the selection. Based on which data attribute the view is currently showing, i.e., `Point Data`, `Cell Data`, or `Row Data`, the selection will select points, cells, or rows, respectively.

Selecting in Line Chart View

`Line Chart View` enables you to select the elements corresponding to the plotted data values. The selection interaction is similar to `Render View`. By default, you are in the interaction mode. You enter selection mode to create a selection by using the buttons in the view toolbar for creating a rectangular selection or a polygonal selection . Once in selection mode, you can click and drag to define the selection region. The selection is created once you release the mouse press.

When a new selection is created, by default, it will clear any existing selection in the view. The selection modifier buttons in the view toolbar can be used to control whether a new selection adds to selected elements , removes points from the selected elements , or toggles it . These modifier buttons are mutually exclusive and modal, i.e., they remain pressed until you click to unpress them or until you press another modifier button. `CTRL` (or `Shift`) and `Alt` can also be used to add to/subtract from the selection.

1.6.3 Creating selections using the `Find Data` panel

Views provide mechanisms to create selections interactively. Selections in chart views and `SpreadSheet View` can be used to select elements with certain data properties, rather than spatial locations ([Fig. 1.82](#)). For a richer data-based selection for selecting elements matching certain criteria, you can use the `Find Data` mechanism in `paraview`.

The `Find Data` panel can be accessed from the `Edit` menu, `:View` menu, or by using the keyboard shortcut `V` or the

The `Find Data` panel can be split into three sections, reflecting how you would use this dialog. The `Create Selection` section helps you define the selection criteria. This identifies which elements, cells or points, are to be selected. The `Selected Data` section shows the results from the most recent selection. They are shown in a tabular view similar to the `Spreadsheet` view. Finally, the `Selection Display` section lets you change how the selected elements are displayed in the active view.

You can create selections `Find Data` panel using the widgets under the `Create Selection` section. First, choose the *data producer*. This is the source or filter from which you want to select elements from. Next choose the *element type*. If you want to select cells, choose `Cell`, for points choose `Point` and so on. The next step is to define the selection criteria. The left-most combo-box is used to select the array of interest. The available options reflect the data array currently available on the dataset. The next combo-box is used to select the operator. Options include the following:

- `is` matches a single value
- `is in range` matches a range of values specified by min and max
- `is one of`` matches a list of comma-separated values
- `is >=` matches all values greater than or equal to the specified value
- `is <=` matches all values lesser than or equal to the specified value

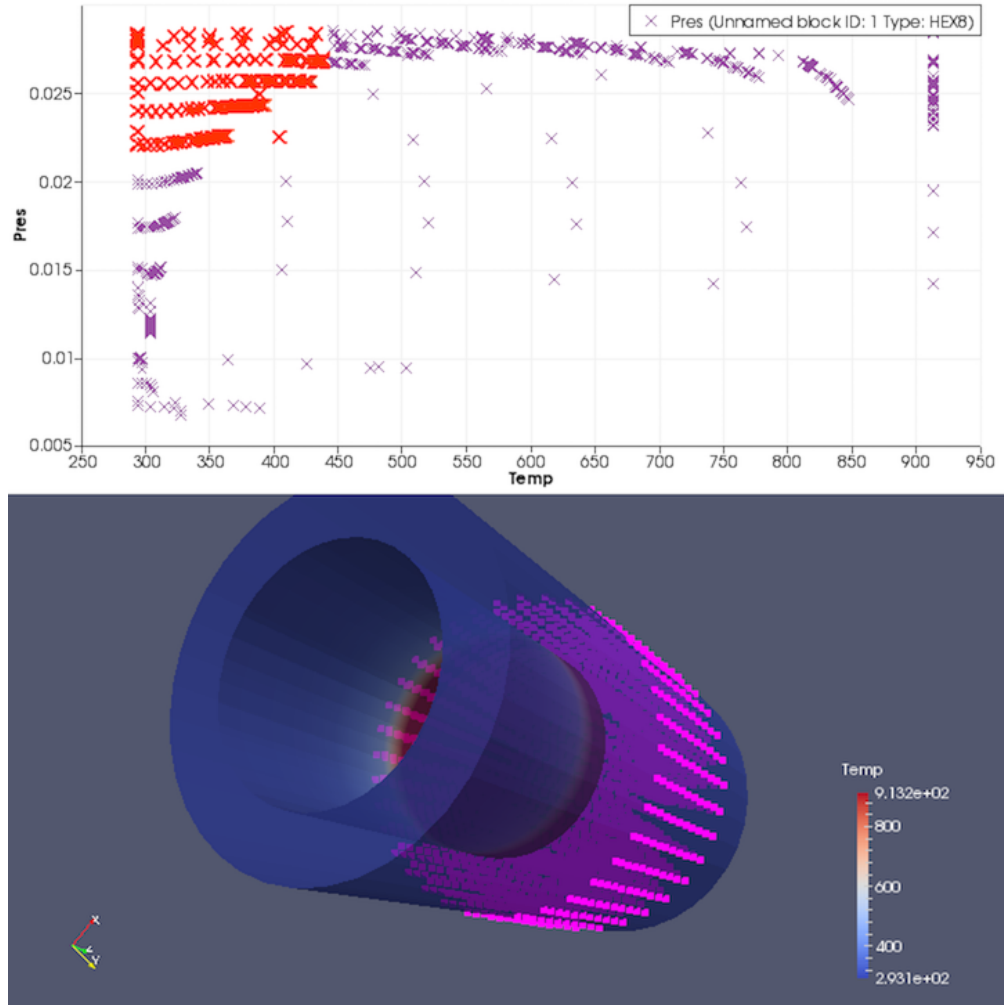


Fig. 1.82: Selection in Line Chart View can be used to locate elements matching attribute criteria. In this visualization, by generating a scatter plot plotting Pres against Temp in the disk_out_ref.ex2 dataset by selecting the top-left corner of the Line Chart View , we can easily locate elements in the high Pres, low Temp regions in the dataset.

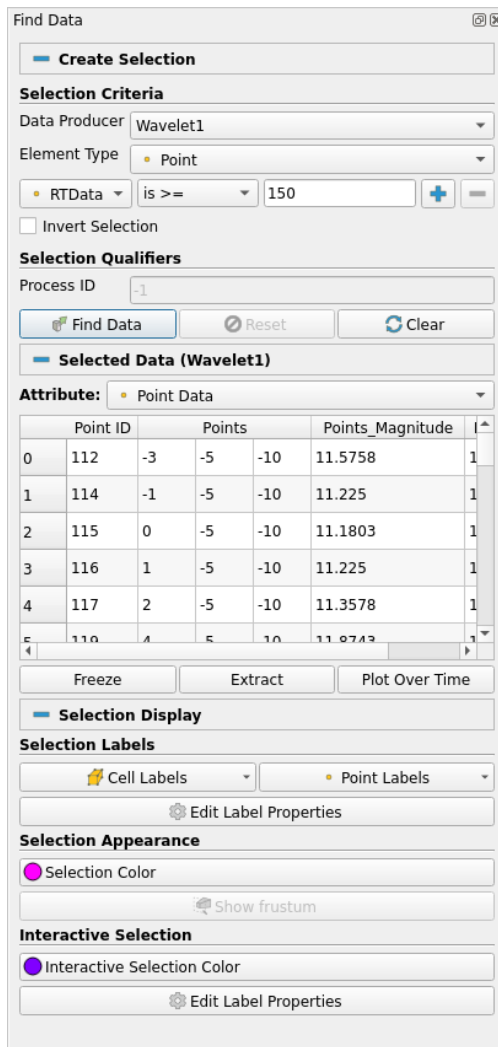


Fig. 1.83: The Find Data panel can be used to find data elements matching specific conditions. In this example, we are selecting all *Points* in *Wavelet1* dataset where *RTData* is ≥ 150 .

- `is min` matches the minimum value for the array for the current time step
- `is max` matches the maximum value for the array for the current time step
- `is <= mean` matches values lesser than or equal to the mean
- `is >= mean` matches values greater than or equal to the mean
- `is mean` matches values equal to the mean within the specified tolerance

Based on your selection of the operator, input widgets will be shown next to this combo-box, where you enter the corresponding values. For example, for `is between`, you enter the min and max values for defining the range in the two text entry widgets.

Multiple selection criteria can be combined together. For example, you want to select all points with `Temp >= 100 and Pres <= mean`, simply setup two expressions using the  button.

Once you are satisfied with the selection criteria, hit the **Find Data** button. On success, the **Current Selection** spreadsheet will update to show the selected elements. Use the **Attribute** combo-box to change which element types are shows in the spreadsheet.

Similar to selecting in views, once you create a selection, any view showing the selected data will also highlight the selected elements, if possible. For example, the **Render View** will show a colored wireframe marking the selected elements, **SpreadSheet View** will highlight the rows, and so on. The **Selection Display** section lets you change how the selection is displayed in the active view. Currently, it is primarily designed for **Render View**. In the future, however, it could support changing selection attributes for other views as well. The available options allow you select the color to use to show the selected elements, as well as the data attributes to use to label the cells/points. For finer control on the label formatting, color, font, etc., use the . That will pop up the **Edit Label Properties** dialog (Fig. 1.84).

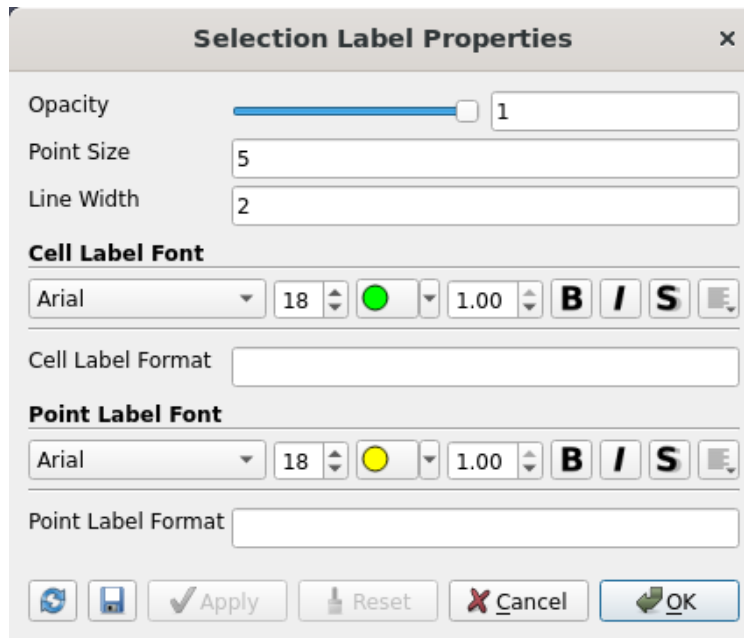


Fig. 1.84: Selection Label Properties dialog for controlling selection labelling parameters.

Did you know?

Besides creating new selections, the **Find Data** dialog can also be used to inspect the current selection made from outside the dialog. For example, if you select elements in the **Render View** using the options described in Section

Section 1.6.2, the Current Selection component in the Find Data dialog will indeed update to reflect the newly selected elements. Furthermore, you can change its display properties and extract this selection using the extraction buttons (which we will cover in Section Section 1.6.6).

1.6.4 Creating selections in Python

Another way to create selections is through ParaView's Python scripting interface. Python functions exist that are analogous to the selection operations available in the ParaView Render View and Find Data dialog. Let's take a look at an example.

```
# import the selection module
from paraview.selection import *

renderView1 = GetActiveView()

# Create an initial rectangular selection in the render view
SelectSurfacePoints(Rectangle=[200, 321, 600, 744], View=renderView1)

# Add points within a polygon in the active view
SelectSurfacePoints(Polygon=[180, 200, 190, 400, 322, 300], Modifier='ADD')

# Subtract points with another rectangle
SelectSurfacePoints(Rectangle=[300, 400, 500, 700], Modifier='SUBTRACT')

# Now extract and show the selected points into another dataset
ExtractSelection()
Show()

# Clear the selection
ClearSelection()
```

The script starts out by importing functions from the `paraview.selection` module. Next, it creates a reference to the active render view and passes it into the selection functions. The first selection function selects points visible in the render view within a rectangular region. The rectangle is defined by bottom left and upper right points, (200, 321) and (600, 744), given in pixel coordinates.

The second selection is of visible points within a polygon defined by the points (180, 200), (190, 400), and (322, 300). In this call, the selection function modifies the existing selection so that newly selected points are added to the selection. This is controlled with the `Modifier` named function parameter. Other options for the `Modifier` parameter are '`SUBTRACT`' , '`TOGGLE`' , and `None` . When the `Modifier` is set to `None` , the previous selection gets replaced with the new selection. The last call to `SelectSurfacePoints` subtracts points from the current selection, which is the combination of the first two selections.

The last lines in this example script extract the currently selected points from the currently active source and shows them on the screen. Lastly, the selection is cleared with the `ClearSelection` function.

Selections by point or cell ID numbers are also possible, as shown in this example:

```
from paraview.selection import *

# Select cell 1 from all blocks in a multiblock data set on process 0
SelectIDs(IDs=[0, 1], FieldType='CELL')
```

(continues on next page)

(continued from previous page)

```
# Add cell 3 from block 4 on process 0 and cell 5 from block 6 on process 1
# to the selection
SelectCompositeDataIDs(IDs=[4, 0, 3, 6, 1, 5], Modifier='ADD')
```

Finally, selections by query expressions are also possible via the Python selection API. As an example, the following selects cells that have the maximum value for a cell variable named EQPS in the currently active source:

```
from paraview.selection import *

QuerySelect(QueryString='EQPS == max(EQPS)', FieldType='CELL')
```

The complete list of selection functions are briefly described below. For full documentation on these functions, you can invoke the `help` function on any of the functions, e.g., `help(SelectSurfaceCells)`.

- `SelectSurfacePoints` - Select visible points within a rectangular or polygon region.
- `SelectSurfaceCells` - Select visible cells within a rectangular or polygon region.
- `SelectSurfaceBlocks` - Select visible blocks within a rectangular region.
- `SelectPointsThrough` - Select all points within a rectangular region regardless of their visibility.
- `SelectCellsThrough` - Select all cells within a rectangular region regardless of their visibility.
- `SelectGlobalIDs` - Select attributes by global IDs.
- `SelectPedigreeIDs` - Select attributes by Pedigree IDs.
- `SelectIDs` - Select attributes by attribute IDs.
- `SelectCompositeDataIDs` - Select attributes by composite attribute IDs.
- `SelectHierarchicalDataIDs` - Select attributes by hierarchical data IDs.
- `SelectThresholds` - Select attributes in a source by thresholding on values in an associated array.
- `SelectLocation` - Select points by location.
- `QuerySelect` - Selection by query expression.
- `ClearSelection` - Clears the selection on the source passed in as a parameter.

1.6.5 Displaying selections

The `Find Data` panel provides easy access to changing the `Selection Display Properties` for the selection in the active view. The same functionality can also be accessed using the `Selection Display Inspector` panel, which is a dockable panel that can be shown/hidden using the `View > Selection Display Inspector` menu. Currently, selection display properties apply to `Render Views` alone. In the future, this will be expanded to other view types, as appropriate.

The `Current Selection` section in the `Find Data` dialog shows the selected elements in a spreadsheet view. You can also make a regular `SpreadSheet View` do the same by checking the  button in the view toolbar to show only selected elements.

1.6.6 Extracting selections

All the types of selections created through mechanisms discussed so far are transient and primarily used for highlighting data. If you want to do further operations on the selected subset, such as extract the selected elements and then save the result out as a new dataset or apply other filters only on the selected elements, then you need to use one of the extract selection filters. The `Extract Selection` and `Plot Selection Over Time` filters fall in this category of filters.

Extract selection

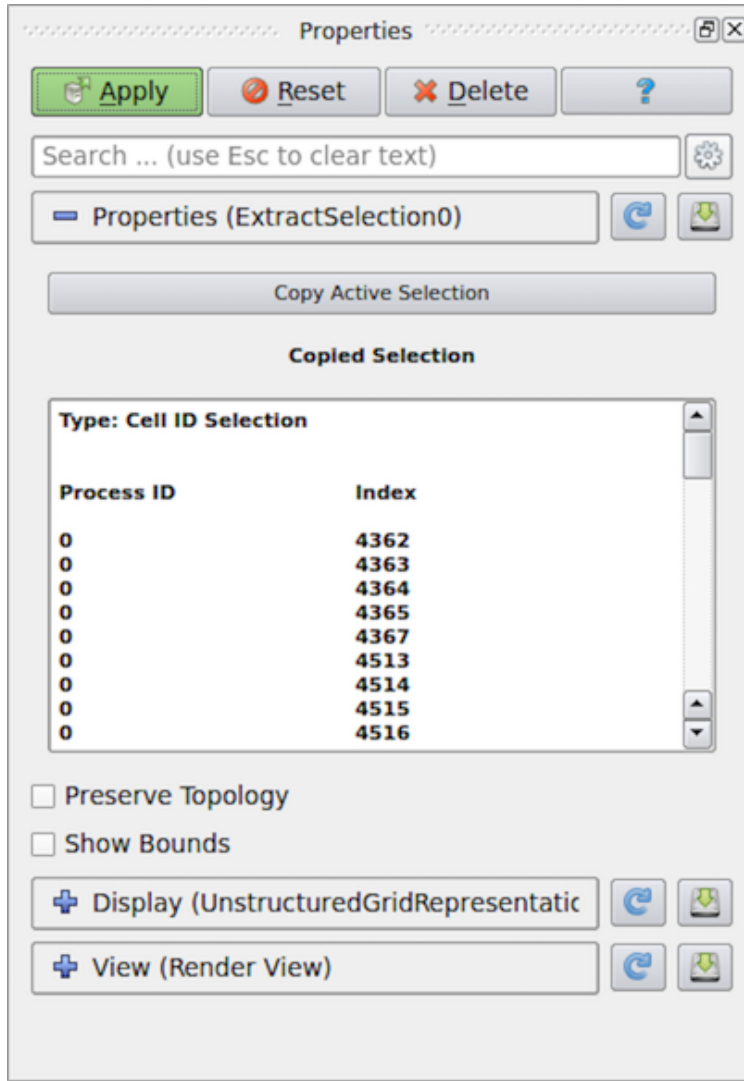


Fig. 1.85: Properties panel showing the properties for the `Extract Selection` filter.)

The `Extract Selection` filter is used to extract the selected elements as a new dataset for further filtering. There are multiple ways of creating this filter. You can use the conventional method for creating filters, i.e., using the `Filters` menu. When the filter is created, if there is any active selection, the filter will automatically copy that selection for convenience. Another way to extract the active selection is using the `Extract Selection` button in the `Find Data` panel (Fig. 1.83).

The `Properties` panel shows what defines the selection. You can update the selection by making a new active selection using any of the mechanisms described earlier in this chapter and then clicking on the `Copy Active Selection` button

on the Properties panel for the Extract Selection filter.

By default, the filter is set up to extract the selected elements alone. This filter also supports passing the entire input dataset through by simply marking which elements are selected. For that, check the Preserve Topology check box on the Properties panel.

Plot selection over time

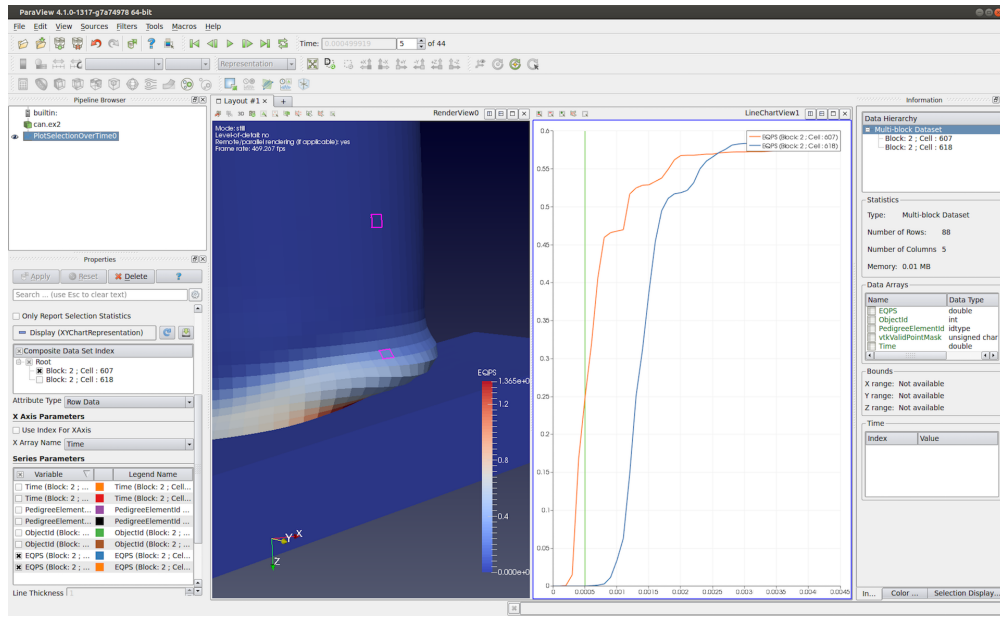


Fig. 1.86: Plot Selection Over Time in action in paraview. The filter provides a convenient way to plot changes in attributes over time for the selected set of cells or for points in a temporal dataset.

Plot Selection Over Time is similar to Extract Selection in the sense that it too extracts the selected elements from the input dataset. However, instead of simply extracting the result, the goal here is to plot attributes at the selected elements over time.

Fig. 1.86 shows an example use of this filter. In this case, we wanted to see how the strain (or *EQPS*) cell attribute changes over time for two specific cells that we selected in the Render View using the view-based selection mechanism. The selected cells are the highlighted elements in the left view. After having selected the cells, we create the Plot Selection Over Time filter using the *Filters > Data Analysis* menu. (You could also use the  from the Data Analysis toolbar.) Similar to the Extract Selection filter, when this filter is created, it copies the active selection. You can change it afterwards using the Copy Active Selection button on the filter's Properties panel. On hitting Apply, paraview will show a visualization similar to the one shown here.

Instead of using the view for defining the selection, you could have used the Find Data panel. In that case, instead of being able to plot each element over time, you will be plotting summaries for the selected subset over time. This is essential since the selected subset can have a varying number of elements over time. The summaries include quantities like minimum, maximum, and median of available variables. You can make the filter always produce these statics alone (even when the selection is created by selecting specific elements in a view) by checking the Only Report Selection Statistics property on the Properties panel for the Plot Selection Over Time filter.

1.6.7 Freezing selections

When extracting selections, you can use views or the **Find Data** panel to define the selection. Since the extraction filters are indeed like any other filters in **ParaView**, they are re-executed any time the input dataset changes, properties on the filter change, or the current time changes. Every time the filter re-executes, it performs the *selection* and *extraction* operations. Thus, if you created the selection using **Render View** to create an id-based selection, the filter will identify which of the elements are of the requested ids and then pass those. For frustum-based selection, it will determine what elements fall within the frustum and extract those. Similarly, with query-based selections created using the **Find Data** panel, the query is re-evaluated. This can result in the selection of different elements with changes in timestep. For example, if you are selecting the cells where the strain is maximum, the selected cell(s) will potentially be different for each time step. Suppose you want to plot the changes in a cell that has maximum strain at the last time step – how can we do that? The answer is using the **Freeze Selection** button on the **Find Data** panel. What that does is convert any type of selection (frustum, query-based) to an id-based selection matching the currently selected element ids. Now you can use this frozen, id-based selection for **Extract Selection** or **Plot Selection Over Time**.

1.7 Animation

In **ParaView**, you can create animations by recording a series of keyframes. At each keyframe, you set values for the properties of the readers, sources, and filters that make up the visualization pipeline, as well as the position and orientation of the camera. Once you have chosen the parameters, you can play through the animation. When you play the animation, you can cache the geometric output of the visualization pipeline in memory. When you replay the animation, playback will be much faster because very little computation must be done to generate the images. Also, the results of the animation can be saved to image files (one image per animation frame) or to a movie file. The geometry rendered at each frame can also be saved in **ParaView**'s PVD file format, which can be loaded back into **ParaView** as a time varying dataset.

1.7.1 Animation View

Animation View is the user interface used to create animations by adding keyframes. It is modeled similar to popular animation and keyframe editing applications with the ability to create tracks for animating multiple parameters. The **Animation View** is accessible from the **View** menu.

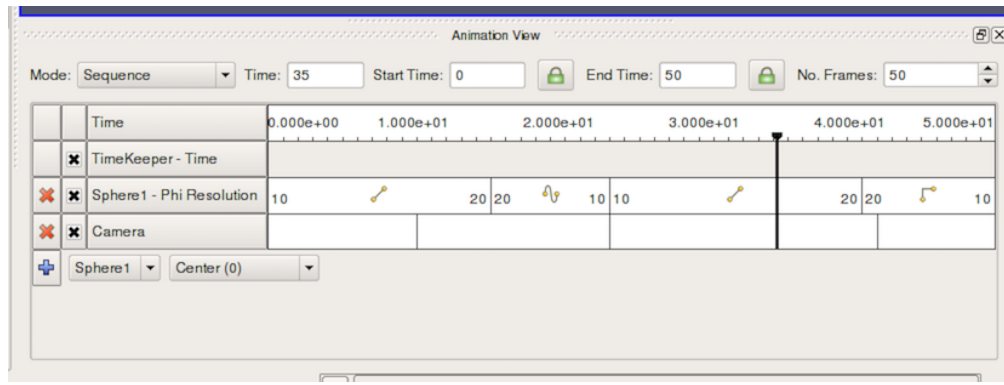


Fig. 1.87: Animation View.

As seen in Fig. 1.87, this view is presented as a table. Above the table are controls that administer how time progresses in the animation. These were discussed briefly in ref{sec:DealingWithTime}. Within the table, the tracks of the animation appear as rows, and animation time is presented as increasing from left-to-right. The first row in the table,

simply labeled Time, shows the total span of time that the animation can cover. The current displayed time is indicated both in the Time field at the top and with a thick, vertical, draggable line within the table.

Along the left side of the Animation View is an expandable list of the names of the animation tracks (i.e., a particular object and property to animate). You choose a data source and then a particular property of the data source in the bottom row. To create an animation track with keyframes for that property, click the + on the left-hand side; this will create a new track. In the figure, tracks already exist for *SphereSource1*'s Phi Resolution property and for the camera's position. To delete a track, press the X button. You can temporarily disable a track by unchecking the check box on the right of the track. To enter values for the property, double-click within the white area to the right of the track name. This will bring up the Animation Keyframes dialog. Double-clicking in the camera entry brings up a dialog like the one in Fig. 1.88.

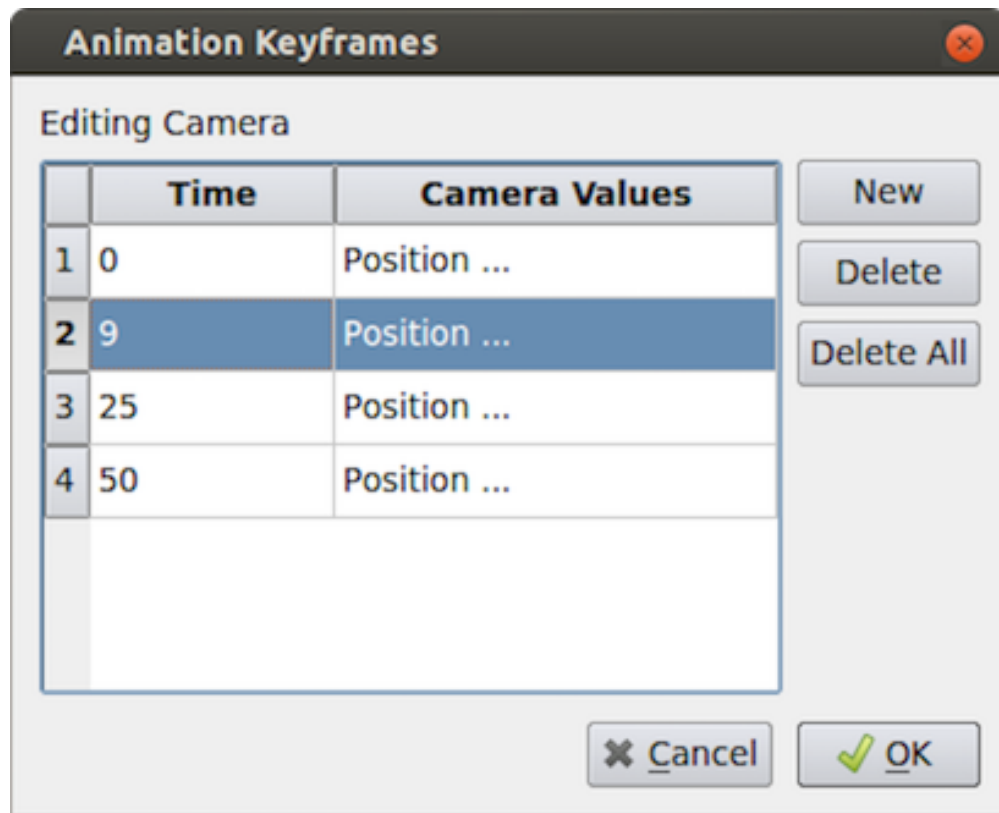


Fig. 1.88: Editing the camera track.

From the Animation Keyframes dialog, you can press New to create new keyframes. You can also press Delete or Delete All to delete some or all of the keyframes. Clicking New will add a new row to the table. In any row, you can click within the Time column to choose a particular time for the keyframe, and you can click in the right-hand column to enter values for the parameter. The exact user interface components that let you set values for the property at the keyframe time vary. When available, you can change the interpolation between two keyframes by double-clicking on the central interpolation column.

Within the tracks of the Animation View, the place in time where each keyframe occurs is shown as a vertical line. The values chosen for the property at that time and the interpolation function used between that value and the next are shown as text, when appropriate. In the previous figure, for example, the sphere resolution begins at 10 and then changes to 20, varying by linear interpolation between them. The camera values are too lengthy to show as text so they are not displayed in the track, but we can easily see that there are four keyframes spaced throughout the animation. The vertical lines in the tracks themselves may be dragged, so you can easily adjust the time at which each keyframe occurs.

Did you know?

You can quickly add a simple animation track and edit the keyframe without using the **Animation View** by using **Animation Shortcut**. First, enable **Show Animation Shortcut** from **Settings** dialog (on the **General** tab, search for the option by name or switch to advanced view). Then, several of the animatable properties on the **Properties** panel will have a  icon. Click this icon to add a new animation track for this property and edit it.



1.7.2 Animation View header

The **Animation View** has a header-bar that lets you control some properties of the animation itself, as you can see in Fig. 1.89.

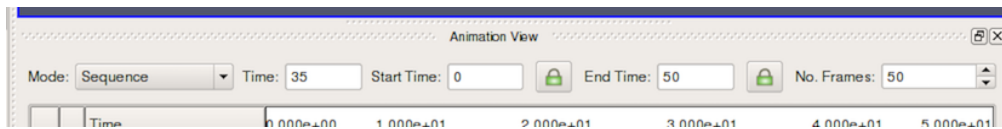


Fig. 1.89: Animation View Header.

Mode controls the animation playback mode. **ParaView** supports three modes for playing animation. In **Sequence** mode, the animation is played as a sequence of images (or frames) generated one after the other and rendered in immediate succession. The number of frames is controlled by the **No. Frames** spinbox at the end of the header. Note that the frames are rendered as fast as possible. Thus, the viewing frame rate depends on the time needed to generate and render each frame.

In **Real Time** mode, the **Duration** spinbox (replacing the **No. Frames** spinbox) indicates the time in seconds over which the entire animation should run. Each frame is rendered using the current wall clock time in seconds, relative to the start time. The animation runs for nearly the number of seconds specified by the **Duration (secs)** spinbox. In turn, the number of frames actually generated (or rendered) depends on the time to generate (or render) each frame.

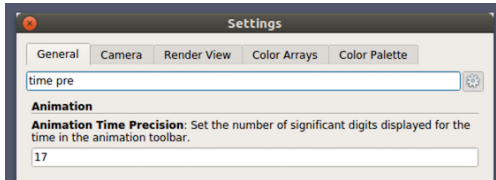
In **Snap To TimeSteps** mode, the number of frames in the animation is determined by the number of time values in the dataset being animated. This is the animation mode used for **ParaView**'s default animations: playing through the time values in a dataset one after the other. Default animations are created by **ParaView** when a dataset with time values is loaded; no action is required to create the animation. Note that using this mode when no time-varying data is loaded will result in no animation at all.

In **Sequence** mode, the final item in the header is the **No. Frames** spinbox. This spinbox lets you pick the total number of frames for the animation. Similarly, in **Real Time** mode, the final line lets you choose the duration of the animation. In **Snap To TimeSteps** mode, the total number of frames is dictated by the dataset and, therefore, the spinbox is disabled.

The **Time** entry-box shows the current animation time, which is the same as shown by a vertical marker in this view. You can change the current animation time by either entering a value in this box, if available, or by dragging the vertical marker. The **Start Time** and **End Time** entry-boxes display the start and end times for the animation. By default, when you load time varying datasets, the start and end times are automatically adjusted to cover the entire time range present in the data. The lock check-buttons to the right of the **Start Time** and **End Time** widgets will prevent this from happening, so that you can ensure that your animation covers a particular time domain of your choosing.

Did you know?

You can change the precision (number of significant digits) displayed by the animation clock by changing the **Animation Time Precision** value under **Settings/Properties Panel Options/Advanced**.



1.7.3 Animating time-varying data

When you load time-varying data, **ParaView** automatically creates a default animation that allows you to play through the temporal domain of the data without manually creating an animation to do so. With the **Animation View**, you can uncouple the data time from the animation time so that you can create keyframes that manipulate the data time during animation as well.

If you double-click in the **TimeKeeper - Time** track, the **Animation Keyframes** dialog, an example of which is shown in [Fig. 1.90](#), appears. In this dialog, you can make data time progress in three fundamentally different ways. If the **Animation Time** radio-button is selected, the data time will be tied to and scaled with the animation time so that, as the animation progresses, you will see the data evolve naturally. If you want to ignore the time varying nature of the data, you can select **Constant Time** instead. In this case, you choose a particular time value at which the data will be displayed for the duration of the animation. Finally, you can select the **Variable Time** radio-button to have full control over data time and to control it as you do any other animatable property in the visualization pipeline. In the example shown in [Fig. 1.90](#), time is made to progress forward for the first 15 frames of the animation, backward for the next 30, and forward for the final 15.

1.7.4 Playing an animation

Once you have designed your animation, you can play through it with the VCR controls toolbar seen in [Fig. 1.15](#).

1.7.5 Animating the camera

Just like you can change parameters on sources and filters in an animation, you can also change the camera parameters. As seen in [Fig. 1.92](#), you can add animation tracks to animate the camera for all the 3D render views in the setup separately. To add a camera animation track for a view, with the view selected, click on the **+** button after choosing **Camera** from the first drop-down menu. The second drop-down list allows you to choose how to animate the camera. There are three possible options, each of which provides different mechanisms to specify the keyframes. It's not possible to change the mode after the animation track has been added, but you can simply delete the track and create a new one.

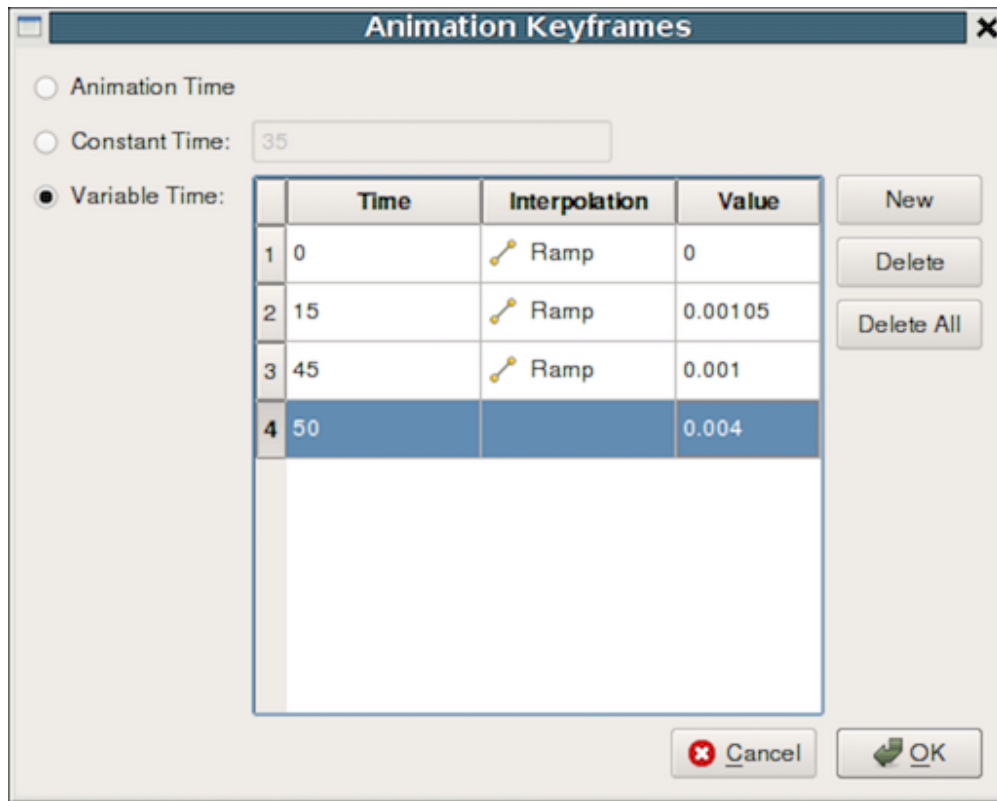


Fig. 1.90: Controlling Data Time with keyframes.

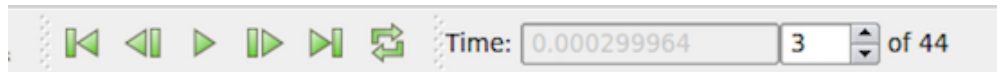


Fig. 1.91: VCR Controls and Current Time Controls toolbars in paraview.

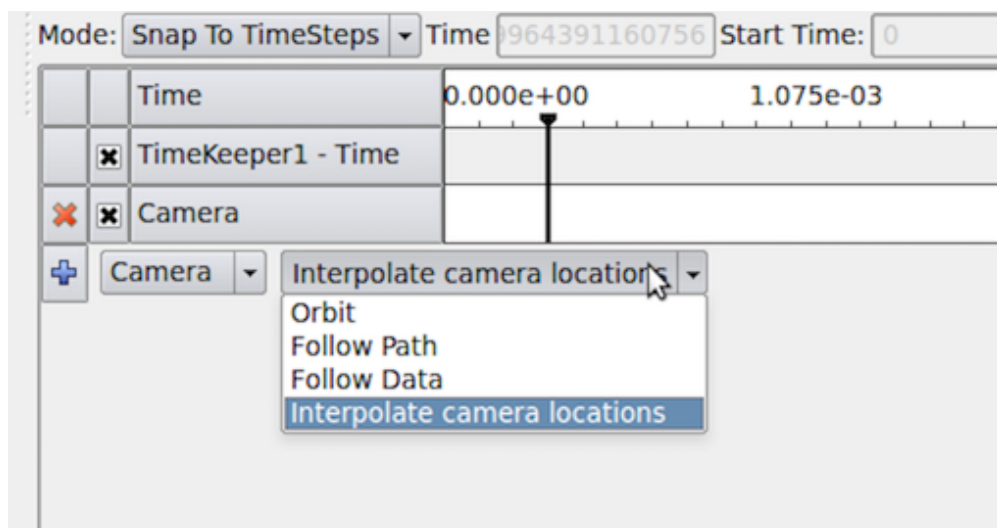


Fig. 1.92: Add camera track.

Interpolate camera locations

In this mode, you specify camera position, focal point, view angle, and up direction at each keyframe. The animation player interpolates between these specified locations. As with other parameters, to edit the keyframes, double-click on the track. It is also possible to capture the current location as a keyframe by using the Use Current button.

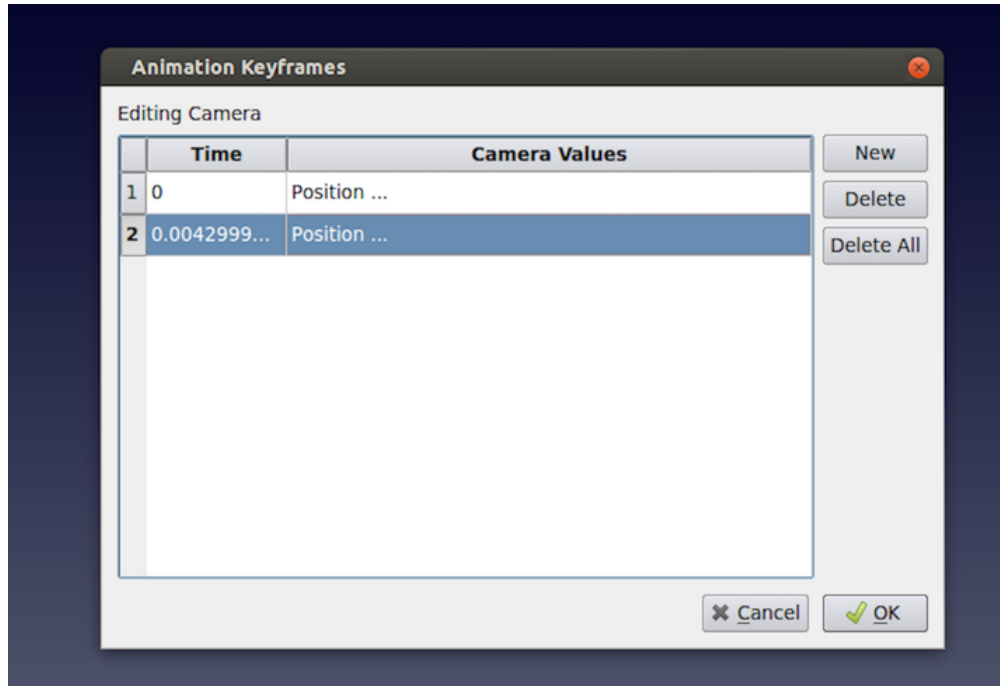


Fig. 1.93: Setting animation parameters.

It can be quite challenging to add keyframes correctly and frequently to ensure that the animation results in a smooth visualization using this mode.

Orbit

This mode makes it possible to quickly create a camera animation in which the camera revolves around objects of interest. Before adding the Camera track, select the objects in the pipeline browser that you want to revolve around; then choose **Orbit** from the Camera combo-box in the Animation View, and hit **+**. This will pop up a dialog where you can edit the orbit parameters such as the center of revolution, the normal for the plane of revolution, and the origin (i.e., a point on the plane where the revolution begins). By default, the **Center** is the center of the bounds of the selected objects, the **Normal** is the current up direction used by the camera, while the origin is the current camera position.

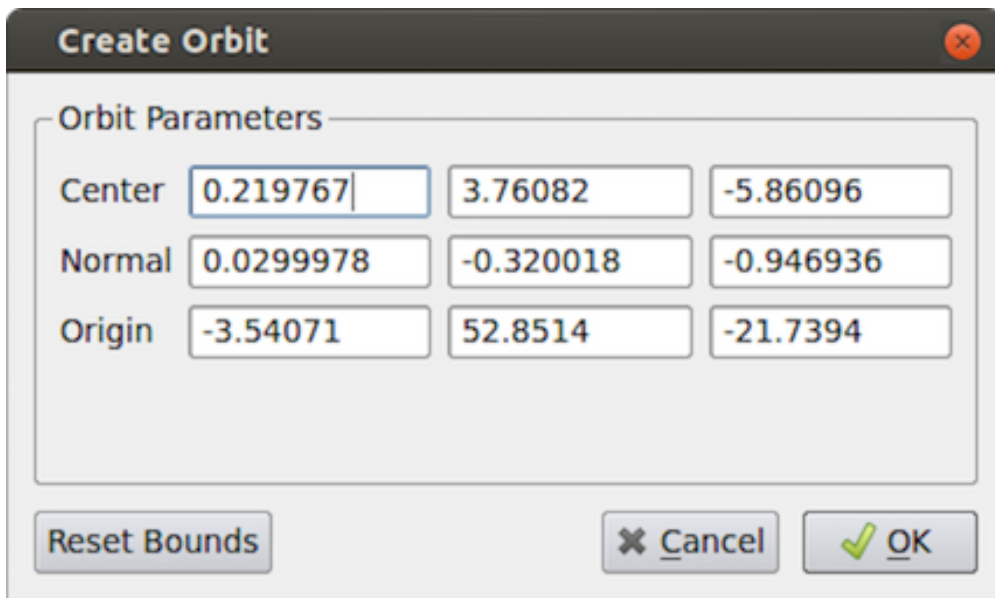


Fig. 1.94: Creating a camera orbit.

Follow path

In this mode, you get the opportunity to specify the path taken by the camera position and the camera focal point. By default, the path is set up to orbit around the selected objects. You can then edit the keyframe to change the paths.

[Fig. 1.95](#) shows the dialog for editing these paths for a keyframe. When Camera Position or Camera Focus is selected, a widget is shown in the 3D view that can be used to set the path. Use `CTRL + Left Click` to insert new control points, and `+ Left Click` to remove control points. You can also toggle when the path should be closed or not.

1.8 Saving Results

In this chapter, we will introduce various ways of saving visualization results in **ParaView**. Results generated throughout the visualization process not only include the images and the rendering results, but also include the datasets generated by filters, the scene representations that will be imported into other rendering applications, and the movies generated from animations.

1.8.1 Saving datasets

You can save the dataset produced by any pipeline module in **ParaView**, including sources, readers, and filters. To save the dataset in `paraview`, begin by selecting the pipeline module in the Pipeline browser to make it the active source. For modules with multiple output ports, select the output port producing the dataset of interest. To save the dataset, use the `File > Save Data` menu or the  button in the Main Controls toolbar. You can also use the keyboard shortcut `CTRL + S` (or `+ S`). The `Save File` dialog ([Fig. 1.96](#)) will allow you to select the filename and the file format. The available list of file formats depends on the type of the dataset you are trying to save.

On accepting a filename and file format to use, `paraview` may show the `Configure Writer` dialog ([Fig. 1.97](#)). This dialog allows you to further customize the writing process. The properties shown in this dialog depend on the selected file format and range from enabling you to `Write All Time Steps`, to selecting the attributes to write in the output file.

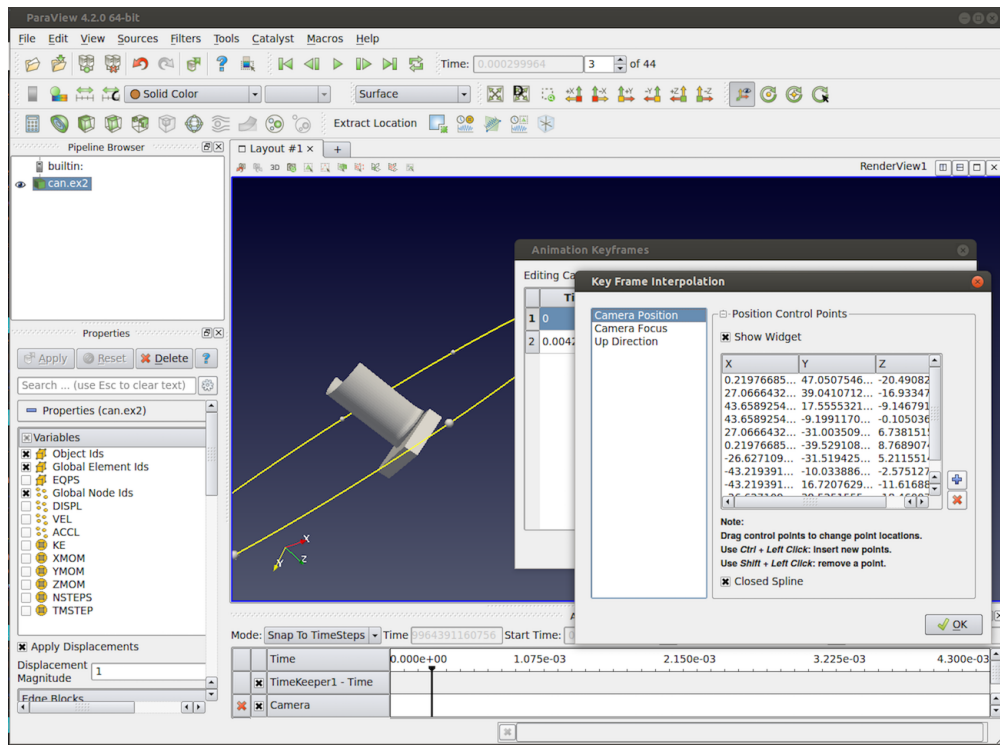


Fig. 1.95: Creating a camera path.

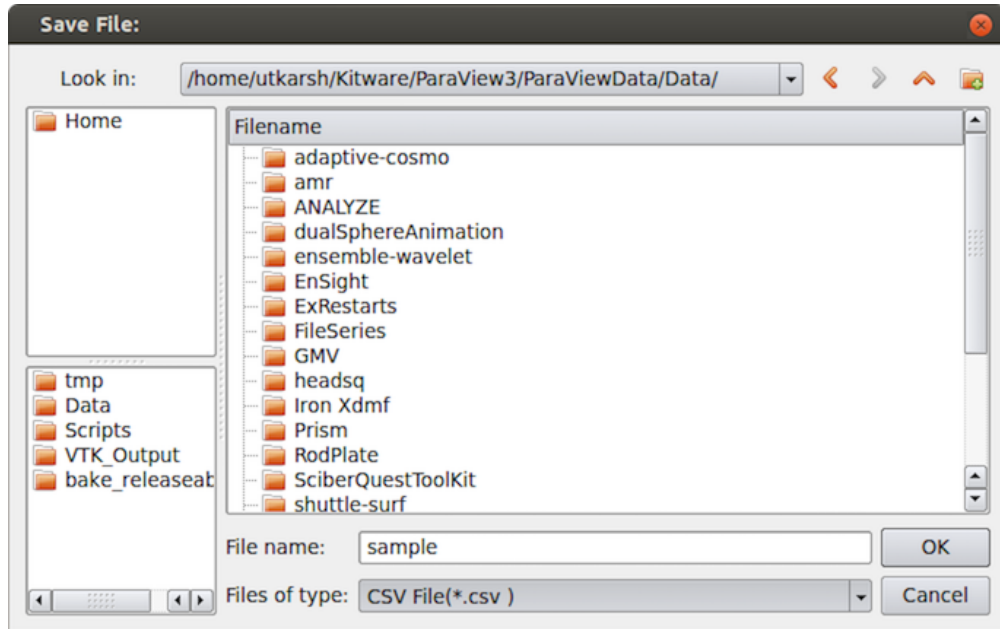


Fig. 1.96: Save File dialog in paraview.

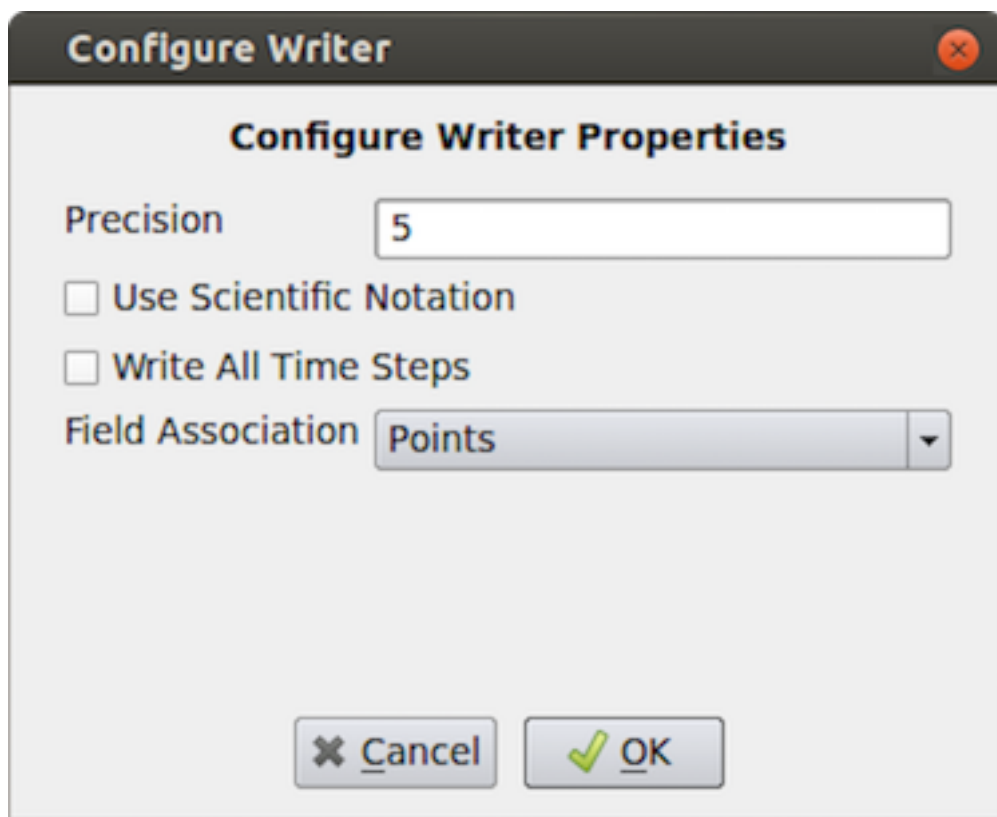


Fig. 1.97: Configure Writer dialog in paraview shown when saving a dataset as a csv file.

In pypython too, you can save the datasets as follows:

```
# Saving the data using the default properties for
# the used writer, if any.
>>> SaveData("sample.csv", source)

# the second argument is optional, and refers to the pipeline module
# to write the data from. If none is specified the active source is used.

# To pass parameters to configure the writer
>>> SaveData("sample.csv", source,
            Precision=2,
            FieldAssociation='Cells')
```

pypython will pick a writer based on the file extension and the dataset type selected for writing, similar to what it does in paraview. Admittedly, it can be tricky to figure out what options are available for the writer. The best way is to use the *Python tracing* capabilities in paraview and to use the generated sample script as a reference ([Section 1.1.6](#)). Make sure you use a similar type of dataset and the same file format as you want to use in your Python script, when tracing, to avoid runtime issues.

1.8.2 Saving rendered results

Views that render results (this includes almost all of the views, except `SpreadSheet View`) support saving images (or screenshots) in one of the standard image formats (PNG, JPEG, TIFF, BMP, PPM). Certain views also support exporting the results in several formats such as PDF, X3D, and VRML.

Saving screenshots

To save the render image from a view in paraview, use the `File > Save Screenshot` menu option. When selected, a file dialog will appear where you can select the file path and format to which the screenshot should be saved. After selecting the image file, the `Save Screenshot Options` dialog ([Fig. 1.98](#)) will be shown. This dialog allows you to select various parameters that controls what image is saved out and how.

If your visualization setup only has 1 view the active tab, then you'll be presented with options shown in ([Fig. 1.98](#)). The available options are as follows.

- **Image Resolution** : This is the target image resolution in pixels. By default, it is set to the current view dimensions. You can change it as needed. If the resolution larger than the current resolution, then ParaView will use *tiling* to render the full image in multiple stages. For reliable results, you may want to present the current aspect ratio. You can use `Tools > Lock View Size Custom` to lock the view size to a suitable aspect ratio.
- **Font Scaling** : When a resolution larger than the current resolution is specified, this allows you to control how the fonts are to be scaled. Default Scale fonts proportionally tries to achieve WYSIWYG as long as the aspect ratio is maintained. This is suitable for saving images targeted for higher DPI (or PPI) display than your screen. Do not scale fonts may be used to avoid font scaling and keep their size in pixels the same as what is currently on the screen. This is suitable for saving images targeted for a larger display with the same pixel resolution.
- **Override Color Palette** : Optionally change the color palette just for saving the screenshot using this dropdown.
- **Stereo Mode** : This option lets you save the image using one of the supported stereo modes.
- **Transparent Background** : If the file format supports it, you can check this option to save the images with a transparent background rather than the current background color.

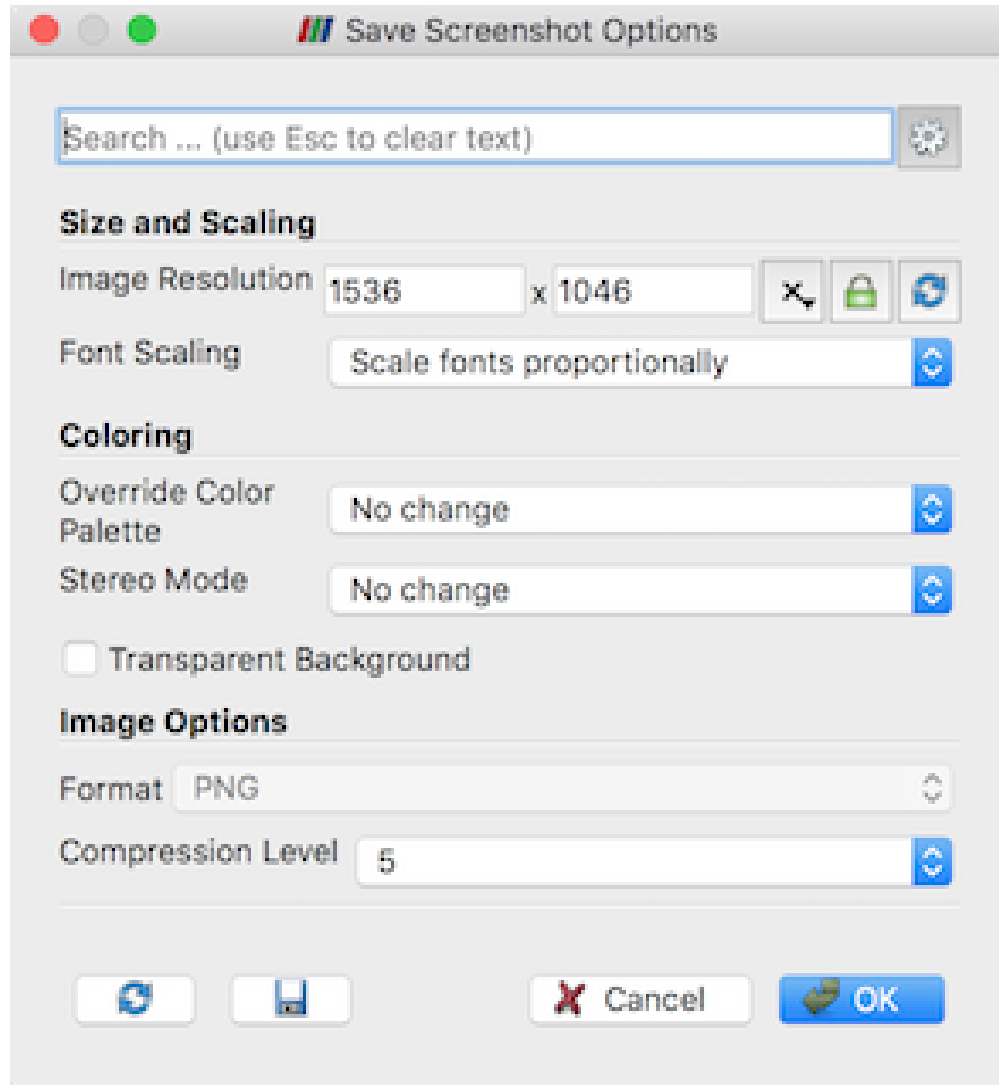


Fig. 1.98: The `Save Screenshot Options` dialog, which is used to customize saving screenshots in paraview.

- **Format** : This shows the file format selected in the file save dialog.

For formats that have different options like compression levels, format-specific options are presented in the Save Screenshot Options dialog. The PNG format has a **Compression Level** option that ranges from 0 (no compression) to 9 (maximum compression). The JPEG format options are **Quality**, which ranges from 0 to 100, and **Progressive**, which enables saving the file as a progressive JPEG. The TIFF file format has a **Compression** option with possible values **None**, **PackBits**, and **Deflate**. The BMP file format has no options.

If the active tab has more than one view, then the Save Screenshot Options dialog has a few more options as shown in Fig. 1.99.

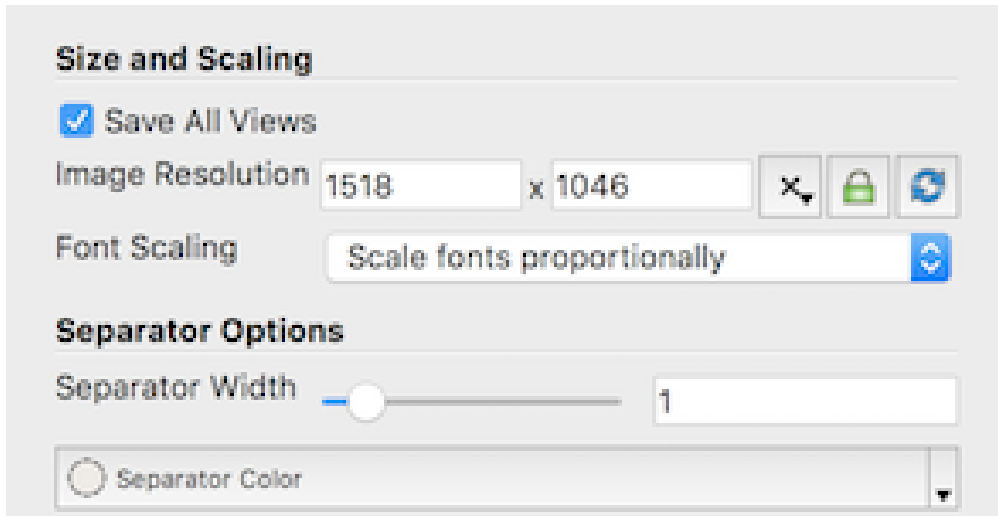


Fig. 1.99: Extra options in Save Screenshot Options dialog available when the active tab has more than 1 view.

- **Save All Views** : Check this to save all the views in the active tab laid out exactly as in the UI. If unchecked, only the active view will be saved.
- **Separator Options** : These control the separator drawn between the views in the generated image. You can specify the **Separator Width** in approximate pixels as well as the **Separator Color**.

To save a screenshot in pypython, you use `SaveScreenshot`.

```
# Save a screenshot from a specific view.
>>> myview = GetActiveView()
>>> SaveScreenshot("aview.png", myview)

# Save all views in a tab
>>> layout = GetLayout()
>>> SaveScreenshot("allviews.png", layout)

# To save a specific target resolution, rather than using the
# the current view (or layout) size, and override the color palette.
>>> SaveScreenshot("aviewResolution.png", myview,
                  ImageResolution=[1500, 1500],
                  OverrideColorPalette="Black Background")
```

As always, you can use *Python tracing* in paraview to trace the exact form of the method to use to save a specific screenshot image.

Exporting scenes

When available, you can export a visualization in a view in several of the supported formats using the *File > Export View* menu option in `paraview`. For a `Render View` (or similar), the available formats include Cinema Database, EPS, PDF, PS, SVG, POV, VRML, WebGL, X3D, and X3DB. On selecting a file as which to export, `paraview` may pop up an `Export Options` dialog that allows you to set up parameters for the exporter, similar to saving datasets (Section 1.8.1).

In addition, from `pvython`, exporting takes the following form (again, just use *Python trace* to figure out the proper form – that's the easiest way).

```
>>> myview = GetActiveView()
>>> ExportView('/tmp/sample.svg', view=myview,
               Plottitle='ParaView GL2PS Export',
               Compressoutputfile=1)
# the arguments after 'view' depend on the exporter selected.
```

1.8.3 Saving animation

To save an animation as a series of images or a video file, you use the *File > Save Animation* menu option. This pops up a file save dialog where you choose where to save the file and which format to use. After selecting the file and format, the `Save Animation Options` dialog (Fig. 1.100) is displayed. This dialog is nearly a clone of the `Save Screenshot Options` dialog (Fig. 1.98), including, optionally, the extra multiview options from Fig. 1.99, with additional format-specific compression options and a few animation-specific parameters. These are as follows:

- **Frame Rate** : When saving the animation as a video file (AVI or Ogg) rather than a series of images, this lets you specify the frame rate for the generated video. It has no effect when saving as a series of images.
- **Frame Window** : If you didn't want to save out the full animation, instead limit to a specific window, you can use this to specify the range of frames to save. If you are generating an animation from a temporal dataset with timesteps, the frame generally corresponds to the timestep number.

On accepting this dialog, you will be able to choose the output file location and format. The available file formats include AVI and Ogg (when available) video formats, as well as image formats such PNG, JPEG, and TIFF. If saving as images, **ParaView** will generate a series of image files sequentially numbered using the frame number as a suffix to the specified filename.

To save animations in `pvython`, you use `SaveAnimation`. The arguments to this function are same as the `SaveScreenshot` with additional parameters for the animation specific options.

```
>>> SaveAnimation('animation.avi', GetActiveView(),
                  FrameWindow = [1, 100],
                  FrameRate = 1)
```

1.8.4 Saving state

Besides saving the results produced by your visualization setup, you can save the state of the visualization pipeline itself, including all the pipeline modules, views, their layout, and their properties. This is referred to as the `Application State`, or just `State`. In `paraview`, you can save the state using the *File > Save State...* menu option. Conversely, to load a saved state file, you can use *File > Load State....*

There are two types of state files that you can save in `paraview`: *ParaView state file (*.pvsm)* and *Python state file (*.py)*. The PVSM files are XML-based text files that are human and machine readable, although not necessarily human friendly for a novice user. However, if you don't plan to read and make sense of the state files, PVSM is the most robust and reliable way to save the application state. For those who want to save the state and then modify it

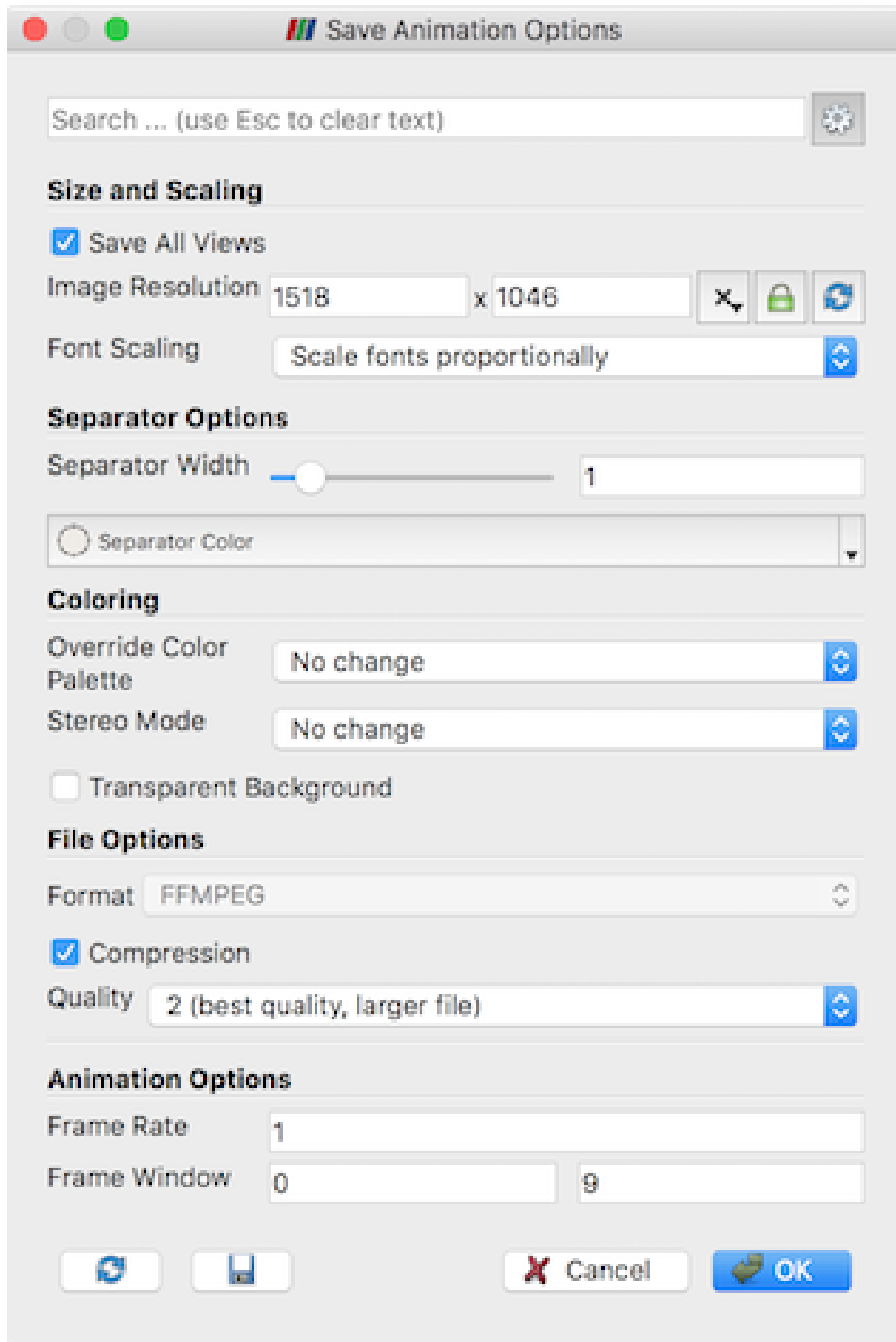


Fig. 1.100: The Save Animation Options dialog in paraview, which is used to customize saving of animation.

manually, using Python state files may be better, as using Python trace simply traces the actions that you perform in the UI as a Python script. Python state files, on the other hand, save the entire current state of the application as a Python script that you can use in paraview or the Python Shell .

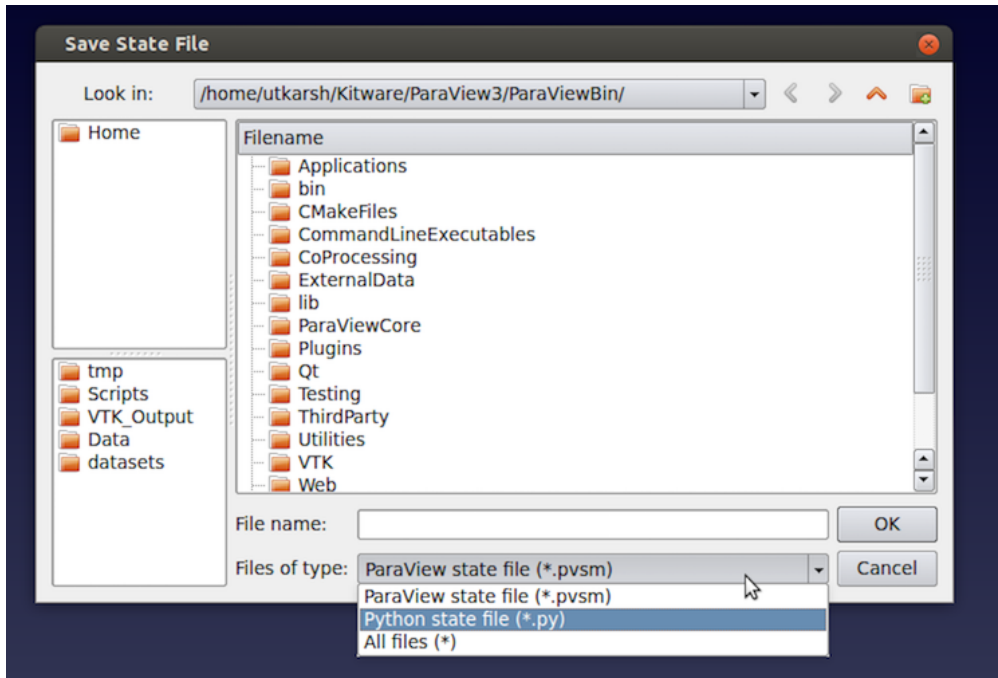


Fig. 1.101: The Save State File dialog in paraview.

To load a state file, you use the *File > Load State...* menu. Note that loading a state file will affect the current visualization state.

If you load a PVSM file this way you will be asked where to search for the data files. There are three available options: **Use File Names From State**, **Search files under specified directory** and **Choose File Names**. If you select **Use File Names From State** then **ParaView** will look for the data at the absolute paths saved in the state file. If you select **Search files under specified directory** then you will see an option to browse for a directory that **ParaView** will search for the files before looking for them in the absolute path in the state file. This defaults to the location of the state file to make sharing state files between computers easier. If you select **Choose File Names** then you will be given a list of file names in the state file and can override each one individually.

You can save/load the PVSM state file in pypython as follows:

```
>>> from paraview.simple import *

# Save the PVSM state file. Currently, this doesn't support
# saving Python state files.
>>> SaveState("sample.pvsm")

# To load a PVSM state file.
>>> LoadState("sample.pvsm")
```

To replace all data files used by state with those under a specific directory, you use the following form:

```
>>> LoadState("sample.pvsm",
            data_directory="[directory path]",
```

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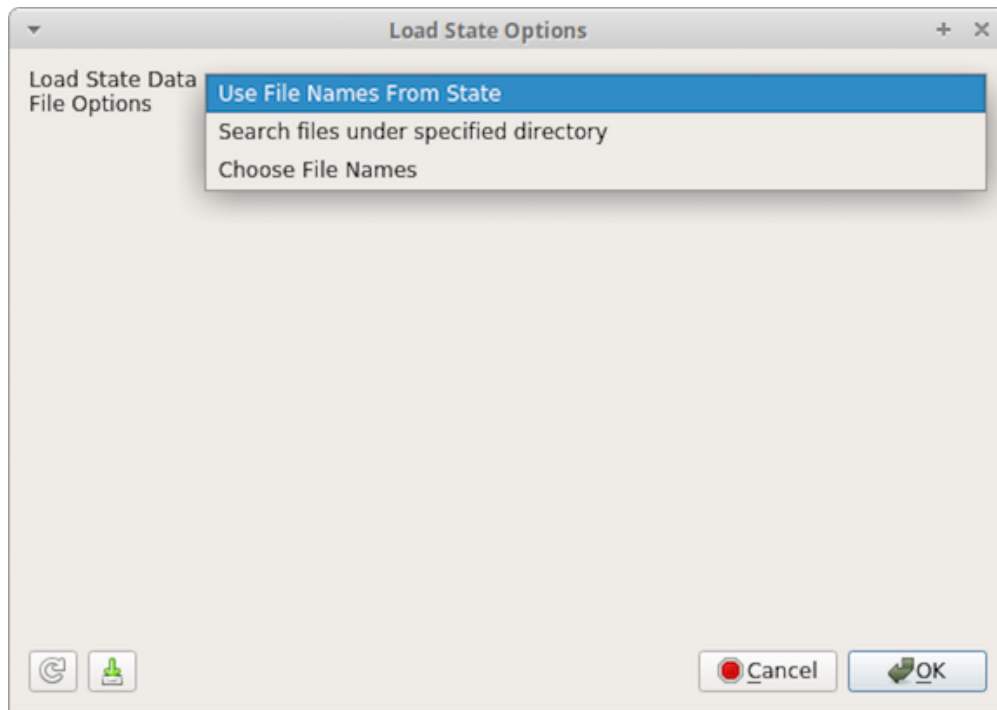


Fig. 1.102: The Load State Options dialog in paraview showing the options for where to find data files.

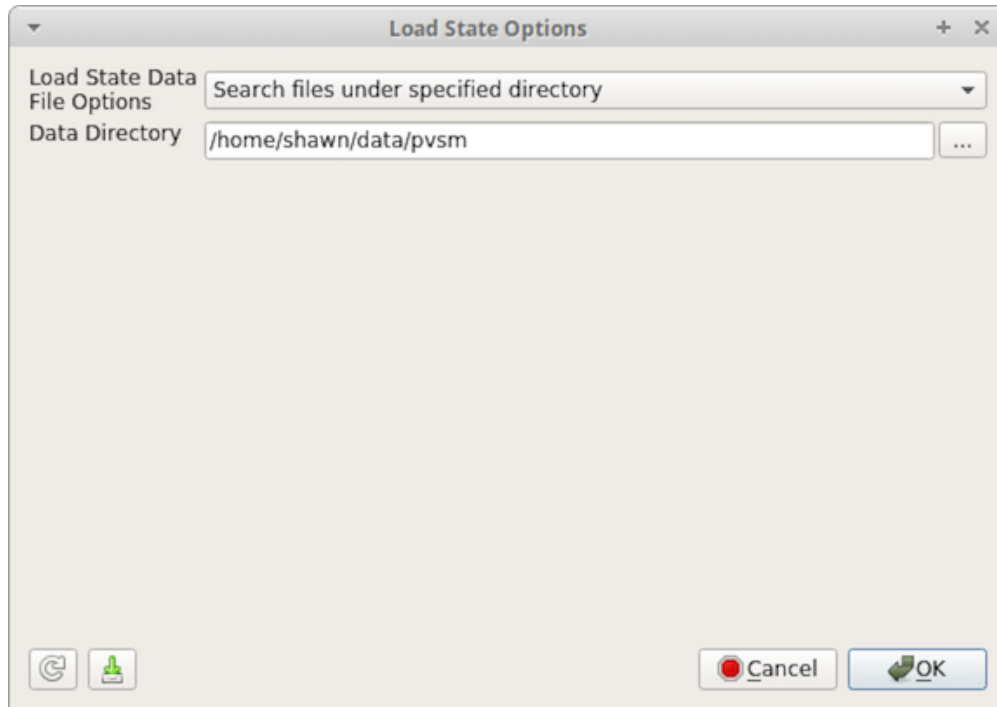


Fig. 1.103: The Load State Options dialog in paraview showing the Search files under specified directory option.

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```
# optionally, restrict to specified directory
restrict_to_data_directory=True)
```

The function signature can become a little more complex if you want to explicitly override filenames used in the state file. It may be easier to use the Python trace capabilities to generate the function call for specific state files. It takes the following form:

```
>>> LoadState("sample.pvsm",
    filenames = [
        # a `dict` object for each reader in statefile to update.
        {
            "name": "[reader name as shown in the pipeline browser]",

            # if multiple readers have the same name, 'id' may be used
            # instead of 'name' where the value is "id" used in the
            # state file for this reader.

            # filename properties and their overridden values for this
            # reader, for example:
            "FileName" : "foo.vtk",
        },
        # multiple such `dict`s can be specified.
    ]
)

# here's an example
>>> LoadState(statefile,
    filenames=[

        {
            'name' : 'can.ex2',
            'FileName' : data_dir + 'can.ex2',
        },
        {
            'name' : 'dataset',
            'FileName' : data_dir + 'disk_out_ref.ex2',
        },
        {
            'name' : 'timeseries',
            'FileName' : [ data_dir + 'dualSphereAnimation/dualSphereAnimation_',
                ↵P00T0000.vtp',
                ↵P00T0001.vtp',
                ↵P00T0002.vtp',
                ↵P00T0003.vtp',
                ↵P00T0004.vtp',
                ↵P00T0005.vtp',
                ↵P00T0006.vtp',
            ],
            'FileName' : [
                data_dir + 'dualSphereAnimation/dualSphereAnimation_',
                data_dir + 'dualSphereAnimation/dualSphereAnimation_',
            ],
        }
    ]
)
```

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```

    ↵P00T0007.vtp',
    ↵P00T0008.vtp',
    ↵P00T0009.vtp',
    ↵P00T0010.vtp']
        },
    ])

```

1.8.5 Extractors

Section 1.8.1 and Section 1.8.2 are two ways of saving datasets and images using actions, i.e., you click a button (or in Python, invoke a function) and the results are saved out immediately. If, for example, you now want to generate the results for another timestep, you have to repeat all the actions. One way to avoid this is to put together a Python script to generate the data and image files and then use that as a macro. An easier way is to use `extractors`. Extractors are a type of pipeline module, similar to sources and filters, but behave more like writers. Similar to filters, they have inputs, unlike sources or filters, however, they produce no output that can be consumed by another pipeline module. Instead, when *activated*, they generate files – which we call extracts.

Since they are just another pipeline module, you use similar mechanisms as sources and filters for creating and configuring these. You use the *Extractors* menu to create them. The Pipeline Browser shows all the extractors present in the visualization. You select one of them by clicking on it in the Pipeline Browser at which point the Properties panel will update to show parameters on the selected extractor.

There are two types of extractors: data extractors and image extractors. The former generate files from datasets produced by sources and filters, while the latter save out rendering results from views. When created, a data extractor by default uses the active source as the input (similar to filters) which an image extractor uses the active view instead.

Extractor Properties

You use the Properties panel view and change extractor properties. The available properties can be grouped into two major groups: first are `Trigger` properties which are common to all extractors, and the second are the `Writer` properties which are parameters specific to type of writer the extractor uses.

`Trigger` properties define when the extractor is activated i.e. under what conditions does the extractor produce extracts. Currently, we support time-based controls. You can select the `Start Time Step`, `End Time Step` or the `Frequency` at which to generate the results. `Frequency` is the number of timesteps per activation thus to write every other timestep, set the `Frequency` to 2, to write every 3rd timestep, set it to 3, and so on.

`Writer` properties are specific to the writer. For data extractors, these will be similar to the writer properties shown in the Configure Writer dialog for the writer described in Section Section 1.8.1. For image extractors, they are similar to the Save Screenshot Options dialog described in Section Section 1.8.2. The `Writer` properties also lets you set a `File Name`. This is the file name to use to save the extracts. Since extractors are designed to generate a new extract every time they are activated, the `File Name` supports patterns that let you make the filename unique per activation. `{timestep}` or `{time}` in the filename are replaced by the timestep index and the time value for each activation. You add leading zeros (or other prefixes) to the numbers using a form such as `{timestep:06d}`. There the timestep will be padded with zeros if the number of digits is less than 6. You should not use absolute paths for specifying the filenames here. We will see how to select prefix to store these extracts under in the next section.

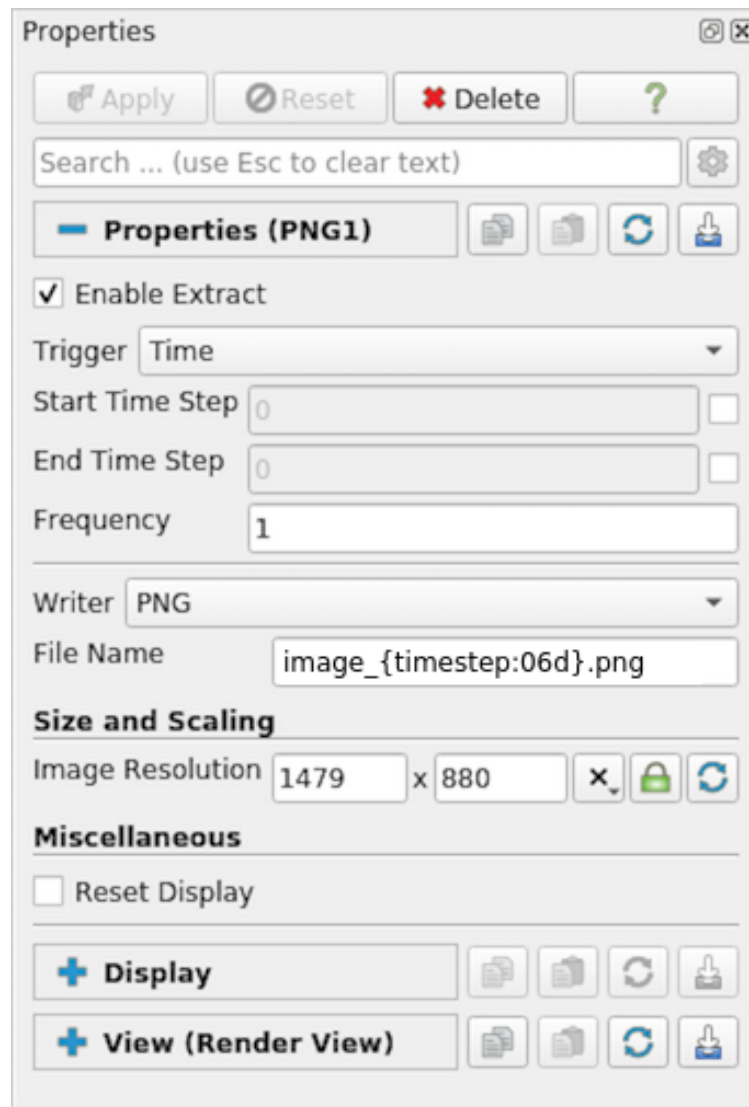


Fig. 1.104: Properties panel showing properties on an image extracts generator for PNG files.

Saving Extracts

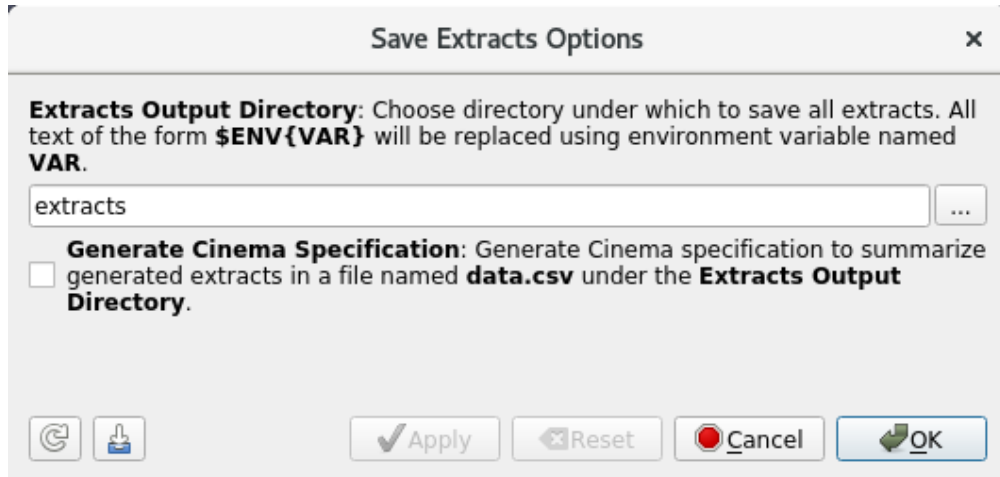


Fig. 1.105: Save Extracts Options dialog shown on *File > Save Extracts....*

Once the extractors have been setup, you can trigger the saving of extract using *File > Save Extracts....* This will pop up the Save Extract Options dialog which lets you configure the extract generation. Extracts Output Directory specifies the root directories under which all extracts are saved. Check Generate Cinema Specification to generate a *data.csv* file under the chosen extracts output directory that summarizes the generated extracts. This can be then used with viewers provided by the Cinema Science project to explore the generated extracts.

Hit Ok and ParaView will animate through all timesteps (similar to using the VCR Controls), activating extractors based on their trigger criteria and then generating extracts. On successful completion you should have files under the chosen root directories.

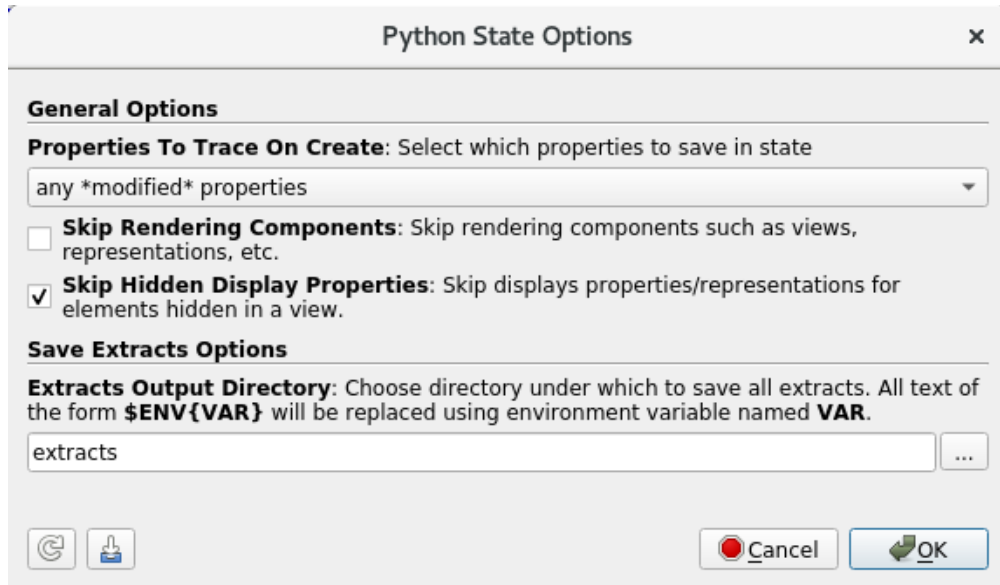


Fig. 1.106: Python State Options dialog shown when saving a Python state file using *File > Save State*.

In addition to generating extracts using the GUI, you can use `pypython` or `pvbash` to generate extract offline. Thus is especially handy for HPC use-cases; you can setup your state using an interactive session and once done save out the state and schedule a non-interactive job for the potentially time-consuming extract generation stage. To do so, setup

your visualization pipeline including the extractors as normal. Then, instead of using `Save Extracts`, use `File > Save State...` and save out a Python state file. The Python State Options dialog has a section similar to `Save Extracts` Options dialog for choosing `Extracts Output Directory` and `Generate Cinema Specification`. Click `Ok` to save the Python script. The Python script has a section near the end of the end as follows:

```
if __name__ == '__main__':
    # generate extracts
    SaveExtracts(ExtractsOutputDirectory='extracts')
```

This is what causes the Python script to save the extracts when the script is executed using `pbatch` or `pvython`.

PARAVIEW REFERENCE MANUAL

2.1 Properties Panel

The **Properties** panel is perhaps the most often used panel in `paraview`. This is the panel you would use to change properties on modules in the visualization pipeline, including sources and filters, to control how they are displayed in views using the **Display** properties, and to customize the view itself. In this chapter, we take a closer look at the **Properties** panel to understand how it works.

2.1.1 Anatomy of the **Properties** panel

Before we start dissecting the **Properties**, remember that the **Properties** panel works on *active* objects, i.e., it shows the properties for the active source and active view, as well as the display properties, if any, for active source in the active view.

Buttons

Fig. 2.1 shows the various parts of the **Properties** panel. At the top is the group of buttons that let you accept, reject your changes to the panel or Delete the active source.

Did you know?

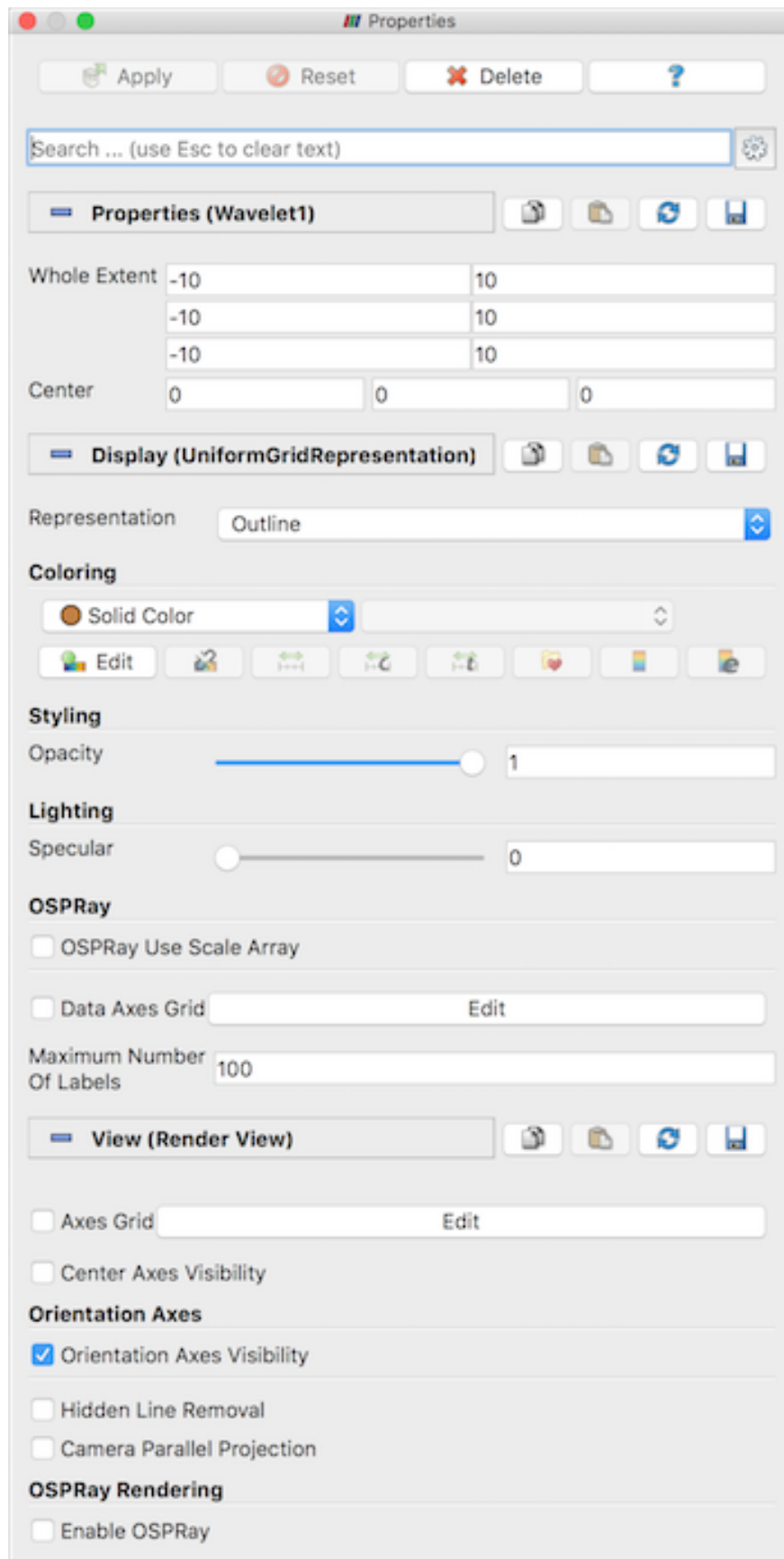
You can delete multiple sources by selecting them using the CTRL (or `⌘`) key when selecting in the **Pipeline Browser** and then clicking on the **Delete** button. Sometimes, the **Delete** button may be disabled. That happens when the selected source(s) have other filters connected to them. You will need to delete those filters first.

Search box

The **Search box** allows you to search for a property by using the name or the label for the property. Simply start typing text in the **Search box**, and the panel will update to show widgets for the properties matching the text.

The **Properties** panel has two modes that control the verbosity of the panel: default and advanced. In the default mode, a smaller subset of the available properties is shown. These are generally the frequently used properties for the pipeline modules. In advanced mode, you can see all the available properties. You can toggle between default and advanced modes using the  button next the **Search box**.

When you start searching for a property by typing text in the **Search box**, irrespective of the current mode of the panel (i.e., default or advanced), all properties that match the search text will be shown.



Did you know?

The Search box is a recurring widget in paraview. Several other panels and dialog boxes, including the **Settings** dialog and the **Color Map Editor**, show a similar widget. Its behavior is also exactly the same as in the case of the **Properties** panel. Don't forget that the  button can be used to toggle between default and advanced modes.

Properties

The **Properties**, **Display**, and **View** sections in the panel show widgets for properties on the active source, its display properties in the active view, and the properties for the active view, respectively. You can collapse/expand each of these sections by clicking on the section name.

To the right of each section name is a set of four buttons. Clicking the  copies the current set of property values to the clipboard while clicking the  will paste those property values into another compatible panel section. Note that the paste icon is enabled only for panel sections where the copied properties can be pasted.

The next two buttons  and  enable customizing the default values used for those properties. Refer to Section 2.11.2 to learn more about customizing default property values.

2.1.2 Customizing the layout

The **Properties** panel, by default, is set up to show the source, display, and view properties on the same panel. You may, however, prefer to have each of these sections in a separate dockable panel. You can indeed do so using the **Settings** dialog accessible from the *Edit > Settings* menu.

On the General tab search of the **properties** panel using the Search box, you should see the setting that lets you pick whether to combine all the sections in one (default), to separate out either the **Display** or **View** sections in a panel, or to create separate panels for each of the sections. You will need to restart paraview for this change to take effect. Also, since the **Apply** and **Reset** buttons only apply to the **Properties** section, they will only be shown in the dock panel that houses it.

2.2 Color maps and transfer functions

One of the first things that any visualization tool user does when opening a new dataset and looking at the mesh is to color the mesh with some scalar data. Color mapping is a common visualization technique that maps data to color, and displays the colors in the rendered image. Of course, to map the data array to colors, we use a transfer function. A transfer function can also be used to map the data array to opacity for rendering translucent surfaces or for volume rendering. This chapter describes the basics of mapping data arrays to color and opacity.

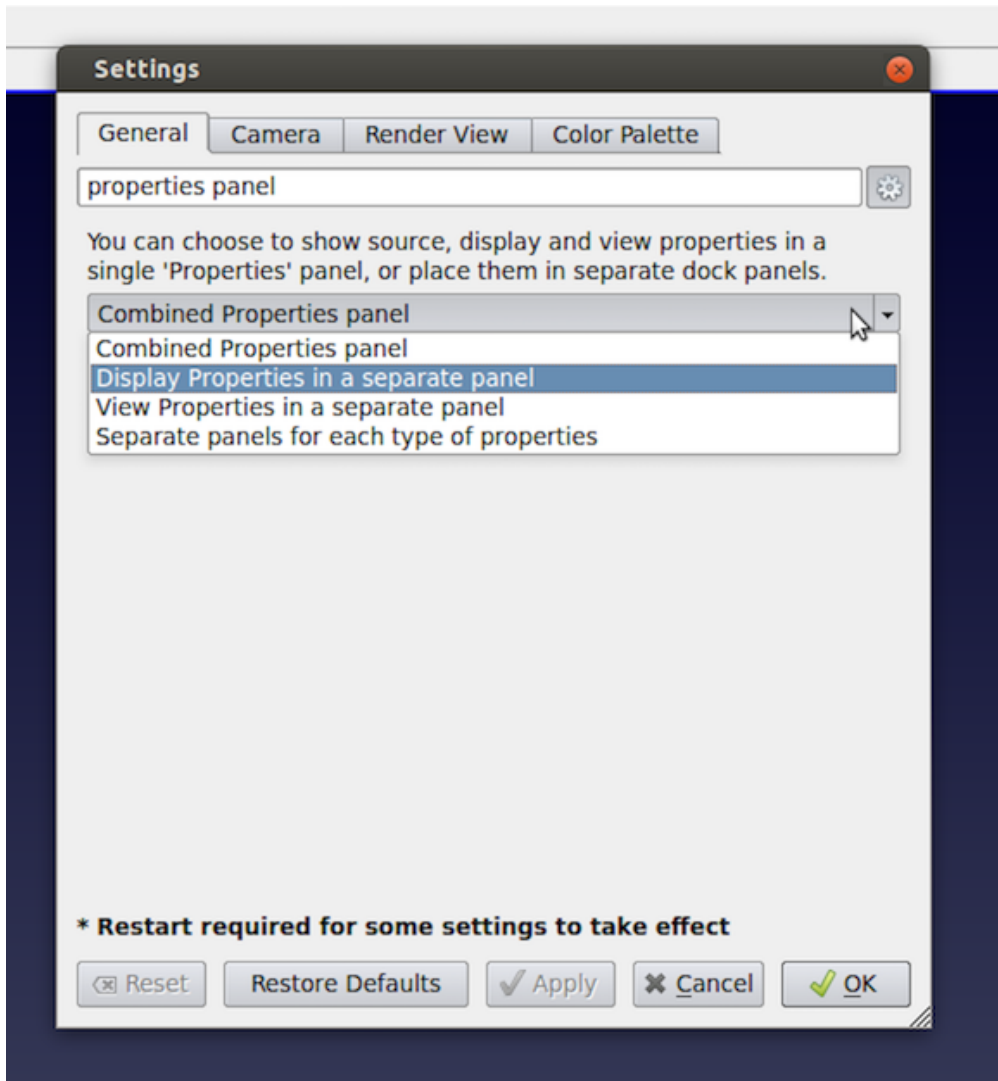


Fig. 2.2: Options for customizing the Properties panel layout using the Settings (left). View properties in a separate dock panel (right).

2.2.1 The basics

Color mapping (which often also includes opacity mapping) goes by various names including scalar mapping and pseudo-coloring. The basic principle entails mapping data arrays to colors when rendering surface meshes or volumes. Since data arrays can have arbitrary values and types, you may want to define to which color a particular data value maps. This mapping is defined using what are called *color maps* or *transfer functions*. Since such mapping from data values to rendering primitives can be defined for not just colors, but opacity values as well, we will use the more generic term *transfer functions*.

Of course, there are cases when your data arrays indeed specify the red-green-blue color values to use when rendering (i.e., not using a transfer function at all). This can be controlled using the `Map Scalars` display property. Refer to Chapter [sec:RenderView:DisplayProperties](#) for details. This chapter relates to cases when `Map Scalars` is enabled, i.e., when the transfer function is being used to map arrays to colors and/or opacity.

In **ParaView**, you can set up a transfer function for each data array for both color and opacity separately. **ParaView** associates a transfer function with the data array identified by its name. The same transfer function is used when coloring with the same array in different 3D views or results from different stages in the pipeline. You can also use [Section 2.2.2](#) to have independent color map by array name and representation.

For arrays with more than one component, such as vectors or tensors, you can specify whether to use the magnitude or a specific component for the color-opacity mapping. Similar to the transfer functions themselves, this selection of how to map a multi-component array to colors is also associated with the array name. Thus, two pipeline modules being colored with the arrays that have the same name will not only be using the same transfer functions for opacity and color, but also the component/magnitude selection.

Common Errors

Beginners find it easy to forget that the transfer function is associated with an array name and, hence, are surprised when changing the transfer function for a dataset being shown in one view affects other views as well. Using different transfer functions for the same variable is discouraged by design in **ParaView**, since it can lead to the misinterpretation of values. If you want to use different transfer functions, despite this caveat, you can use the Separate Color Map feature (see [Section 2.2.2](#)).

There are separate transfer functions for color and opacity. The opacity transfer function is used for volume rendering, and it is optional when used for surface renderings.

Color mapping in paraview

You can pick an array to use for color mapping, using either the **Properties** panel or the **Active Variables Controls** toolbar. You first select the array with which to color and then select the component or magnitude for multi-component arrays. **ParaView** will either use an existing transfer function or create a new one for the selected array.

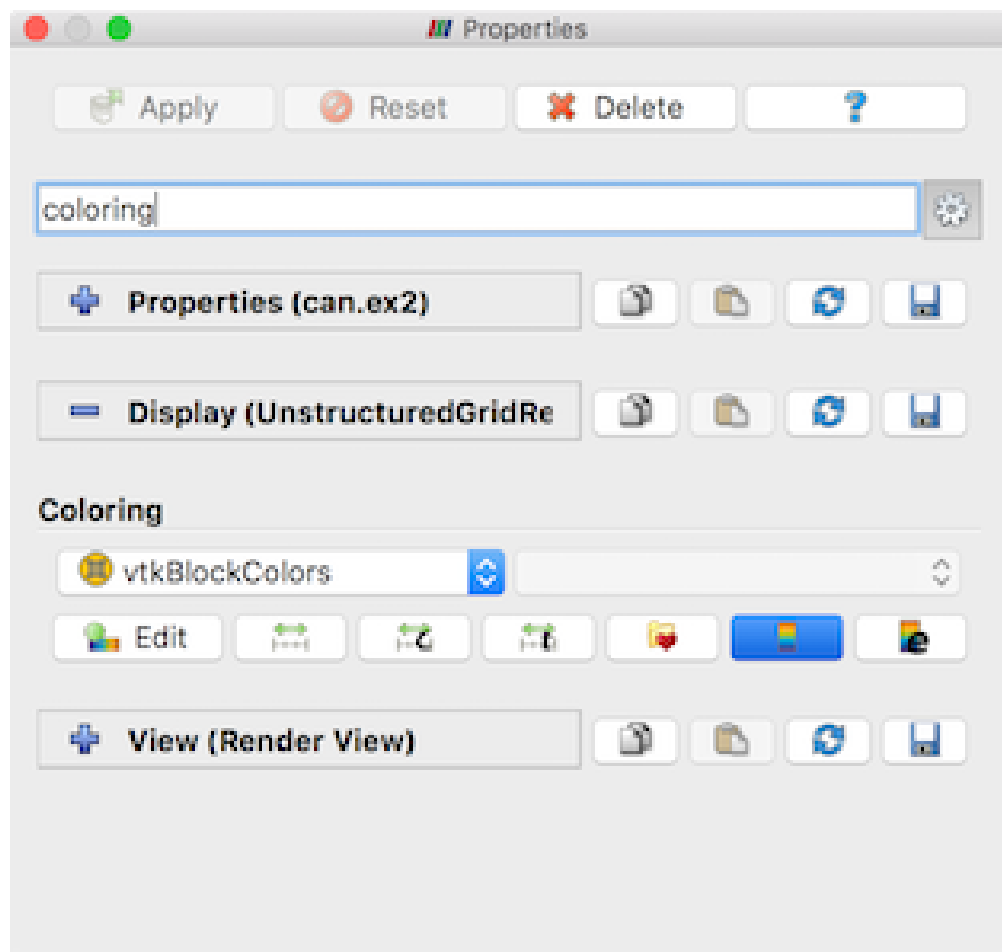
Color mapping in pvpthon

Here's a sample script for coloring using a data array from the `disk_out_ref.ex2` dataset.

```
from paraview.simple import *

# create a new 'ExodusIIReader'
reader = ExodusIIReader(FileName=['disk_out_ref.ex2'])
reader.PointVariables = ['V']
reader.ElementBlocks = ['Unnamed block ID: 1 Type: HEX8']
```

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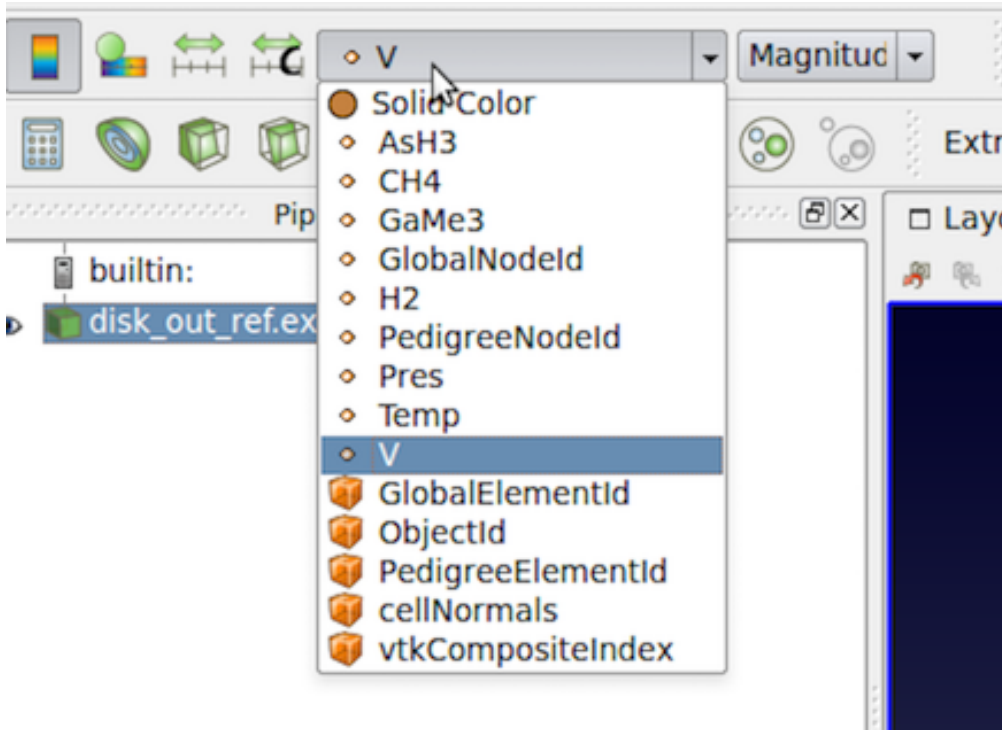


Fig. 2.3: The controls used for selecting the array to color within the `Properties` panel (top) and the `Active Variables Controls` toolbar (bottom).

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```
# show data in view
display = Show(reader)

# set scalar coloring
ColorBy(display, ('POINTS', 'V'))

# rescale color and/or opacity maps used to include current data range
display.RescaleTransferFunctionToDataRange(True)
```

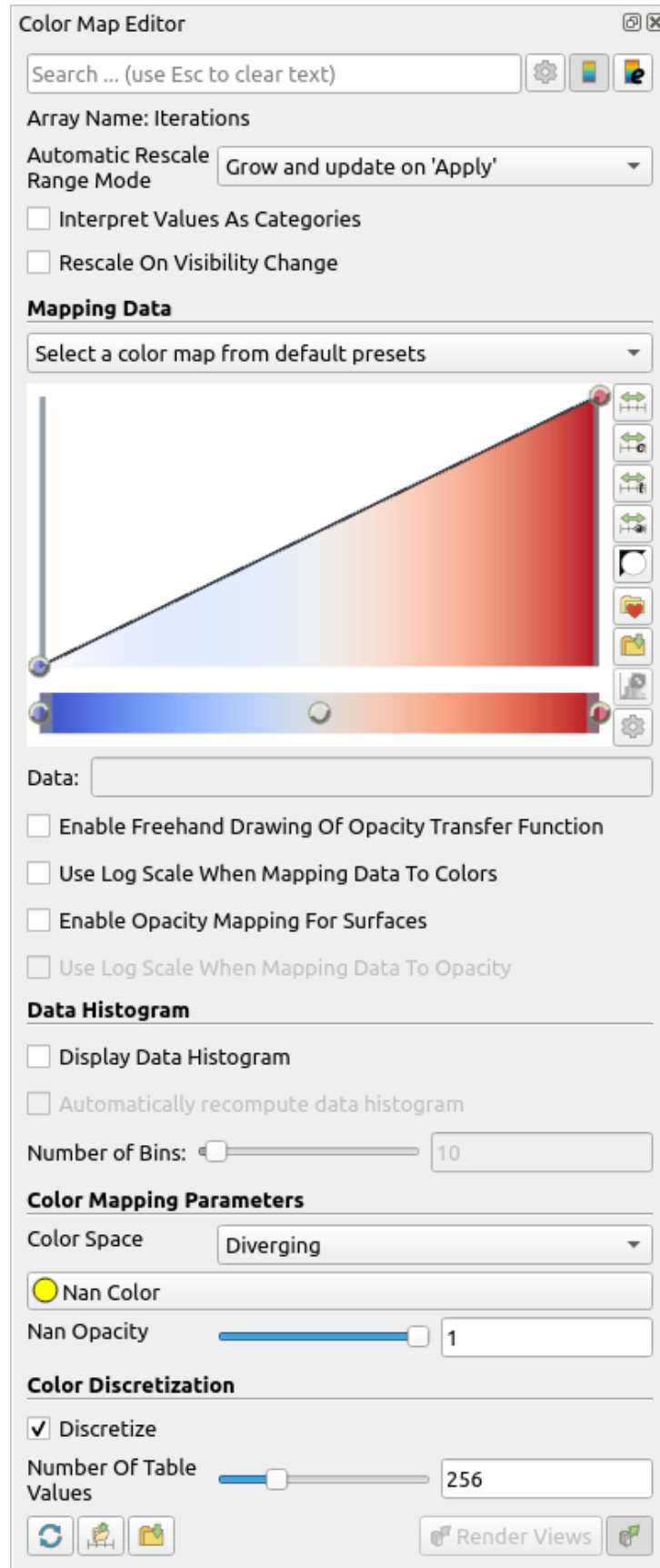
The `ColorBy` function provided by the `simple` module ensures that the color and opacity transfer functions are set up correctly for the selected array, which is using an existing one already associated with the array name or is creating a new one. Passing `None` as the second argument to `ColorBy` will display scalar coloring.

2.2.2 Editing the transfer functions in paraview

In `paraview`, you use the `Color Map Editor` to customize the color and opacity transfer functions. You can toggle the `Color Map Editor` visibility using the `View > Color Map Editor` menu option.

As shown in Fig. 2.4, the panel follows a layout similar to the `Properties` panel. The panel shows the properties for the transfer function, if any, used for coloring the active data source (or filter) in the active view. If the active source is not visible in the active view, or is not employing scalar coloring, then the panel will be empty.

Similar to the `Properties` panel, by default, the commonly used properties are shown. You can toggle the visibility of advanced properties by using the button. Additionally, you can search for a particular property by typing its name



in the Search box.

Whenever the transfer function is changed, we need to re-render, which may be time consuming. By default, the panel requests a render on every change. To avoid this, you can toggle the button. When unchecked, you will need to manually update the panel using the Render Views button.

The button restores the application default settings for the current color map.

The and buttons save the current color and opacity transfer function, with all its properties, as the default transfer function. ParaView will use it next time it needs to set up a transfer function to color a new data array. The button saves the transfer function as default for an array of the same name while the button saves the transfer function as default for all arrays. Note that this will not affect transfer functions already setup. Also this is saved across sessions, so ParaView will remember this even after restart.

Separate Color Map



Fig. 2.5: The Separate Color Map button

In order to force ParaView to use a separate color map on the current Active Representation, click on the button shown in Fig. 2.5. A separate color map is not shared across representations by name, but is instead uniquely associated with the array name and the representation.

This can also easily be done in Python:

```
from paraview.simple import *

Wavelet()
wavelet1Display = Show()
wavelet1Display.SetRepresentationType('Surface')

# set scalar coloring
ColorBy(wavelet1Display, 'RTData')

# set the usage of a Separate Color Map
wavelet1Display.UseSeparateColorMap = True

# or use the ColorBy interface directly
ColorBy(wavelet1Display, 'RTData', separate = True)

# display the same data in another view for comparison with different color map
# get layout
layout1 = GetLayout()

# split cell
layout1.SplitHorizontal(0, 0.5)

renderView1 = GetActiveView()
```

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```

# Create a new 'Render View'
renderView2 = CreateView('RenderView')

# place view in the layout
layout1.AssignView(2, renderView2)

# set active view
SetActiveView(renderView2)

wavelet2Display = Show()
wavelet2Display.SetRepresentationType('Surface')

# Use the ColorBy interface to create a separated color map
ColorBy(wavelet2Display, 'RTData', separate = True)

# get separate color transfer function/color map for 'RTData'
separate_wavelet2Display_RTDataLUT = GetColorTransferFunction('RTData', wavelet2Display,
    ↴separate=True)

# Apply a preset using its name.
separate_wavelet2Display_RTDataLUT.ApplyPreset('Cold and Hot', True)

ResetCamera(renderView1)
ResetCamera(renderView2)
RenderAllViews()

```

Mapping data

The **Mapping Data** group of properties controls how the data is mapped to colors or opacity. The transfer function editor widgets are used to control the transfer function for color and opacity. The panel always shows both the transfer functions. Whether the opacity transfer function gets used depends on several things:

- When doing surface mesh rendering, it will be used only if **Enable opacity mapping for surfaces** is checked
- When doing volume rendering, the opacity mapping will always be used.

To map the data to color using a log scale, rather than a linear scale, check the **Use log scale when mapping data to colors**. It is assumed that the data is in the non-zero, positive range. **ParaView** will report errors and try to automatically fix the range if it is ever invalid for log mapping.

The range of a color map is a very important property that controls the mapping of data values to colors. The range can be automatically updated in a number of situations for convenience. How the range is updated is controlled by the **Automatic Rescale Range Mode** property in the **Color Map Editor**. When **Never** is selected, the data range will never be updated automatically. When **Grow and update on 'Apply'** is selected, **ParaView** will grow the color-opacity map range to include the current data range every time you hit **Apply** on the **Properties** panel. Thus, when the data range changes, if the timestep is changed, the color-opacity map range won't be affected. To grow the range on change in timestep as well, use the **Grow and update every timestep** option. Now the range will be updated on **Apply** as well as when the timestep changes. **Grow** indicates that the color-opacity map range will only be increased, never shrunk, to include the current data range. If you want the range to match the current data range exactly, then you should use the **Clamp and update every timestep** option. Now the range will be clamped to the exact data range each time you hit **Apply** on the **Properties** panel or when the timestep changes. The initial value for the **Automatic Rescale Range Mode** is controlled by the General setting **Transfer Function Reset Mode** in the

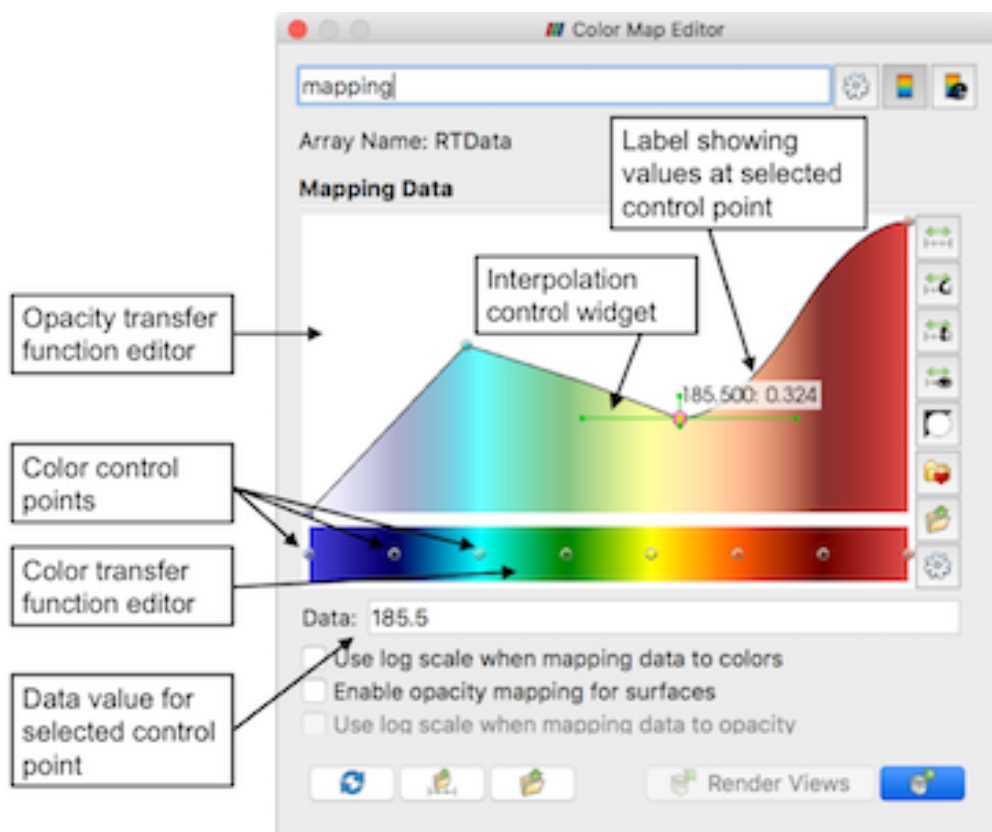


Fig. 2.6: Transfer function editor and related properties

Settings dialog (see [Section 2.11.1](#)).

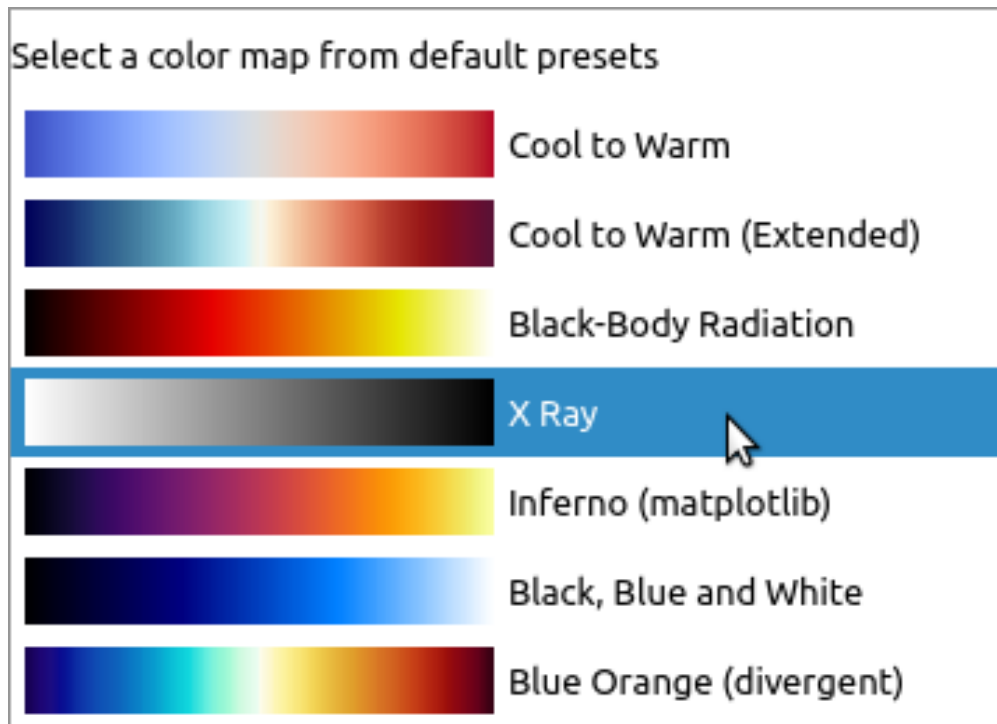
Transfer function editor

Using the transfer function editors is pretty straightforward. Control points in the opacity editor widget and the color editor widget are independent of each other. To select a control point, click on it. When selected, the control point is highlighted with a red circle and data value associated with the control point is shown in the Data input box under the widget. Clicking in an empty area will add a control point at that location. To move a control point, click on the control point and drag it. You can fine tune the data value associated with the selected control point using the Data input box. To delete a control point, select the control point and then type the Delete key. Note that the mouse pointer should be within the transfer function widget for Qt to send the event to the editor widget correctly. While the end control points cannot be moved or deleted, you can rescale the entire transfer function to move the control points, as is explained later.

In the opacity transfer function widget, you can move the control points vertically to control the opacity value associated with that control point. In the color transfer function widget, you can double click on the control point to pop up a color chooser widget to set the color associated with that control point.

The opacity transfer function widget also offers some control over the interpolation between the control points. Double click on a control point to show the interpolation control widget, which allows for changing the sharpness and midpoint that affect the interpolation. Click and drag the control points to see the change in interpolation.

The combo box at the top of the transfer function editor is used to quickly switch between the “Default” presets. Which presets are default ones can be configured from the [Color Preset manager](#), which can be accessed with the  described later.



The several control buttons on the right side of the transfer function widgets support the following actions:

-  : Rescales the color and opacity transfer functions using the data range from the data source selected in the Pipeline browser, i.e., the active source. This rescales the entire transfer function. Thus, all control points including the intermediate ones are proportionally adjusted to fit the new range.

- : Rescales the color and opacity transfer functions using a range provided by the user. A dialog will be popped up for the user to enter the custom range.
- : Rescales the color and opacity transfer functions to the range of values for data over all timesteps. This operation may be costly as data for all timesteps needs to be read.
- : Rescales the color and opacity transfer functions using the range of values for the elements (cells or points) visible in the view. This operation assigns the entire range of colors to visible elements which may reveal patterns not visible otherwise.
- : Inverts the color transfer function by moving the control points, e.g., a red-to-green transfer function will be inverted to a green-to-red one. This only affects the color transfer function and leaves the opacity transfer function untouched.
- : Loads the color transfer function from a preset. The **Color Preset** manager dialog pops up to enable you to choose one of the color maps included with ParaView or import presets from a file.
- : Saves the current color transfer function to presets. The **Color Preset** manager dialog pops up to let you name the transfer function and export the transfer function to a file. The opacity function can also be saved with the transfer function. The preset will be added under the **Default** and **User** groups.
- : This toggles the detailed view for the transfer function control points. This is useful to manually enter values for the control points rather than using the UI.

Color mapping parameters

The **Color Mapping Parameters** group of properties provides additional control over the color transfer function, including control over the color interpolation space, which is either RGB, HSV, Lab, Diverging, or Lab/CIEDE2000. To color data values falling below or above the range of the color map with special colors, enable the advanced **Use Below Range Color** and **Use Above Range Color** options, respectively. You can choose different colors for data falling on either side of the range. When color mapping floating point arrays with NaNs, you can select the color and opacity to use for NaN values. You can also affect whether the color transfer function uses smooth interpolation or discretizes the map into a fixed number of colors.

2.2.3 Editing the transfer functions in pypython

In pypython, you control the transfer functions by getting access to the transfer function objects and then changing properties on those. The following script shows how you can access transfer functions objects.

```
from paraview.simple import *

# You can access the color and opacity transfer functions
# for a particular array as follows. These functions will
# create new transfer functions if none exist.
# The argument is the array name used to locate the transfer
# functions.
>>> colorMap = GetColorTransferFunction('Temp')
>>> opacityMap = GetOpacityTransferFunction('Temp')
```

Once you have access to the color and opacity transfer functions, you can change properties on these similar to other sources, views, etc. Using the Python tracing capabilities to discover this API is highly recommended.

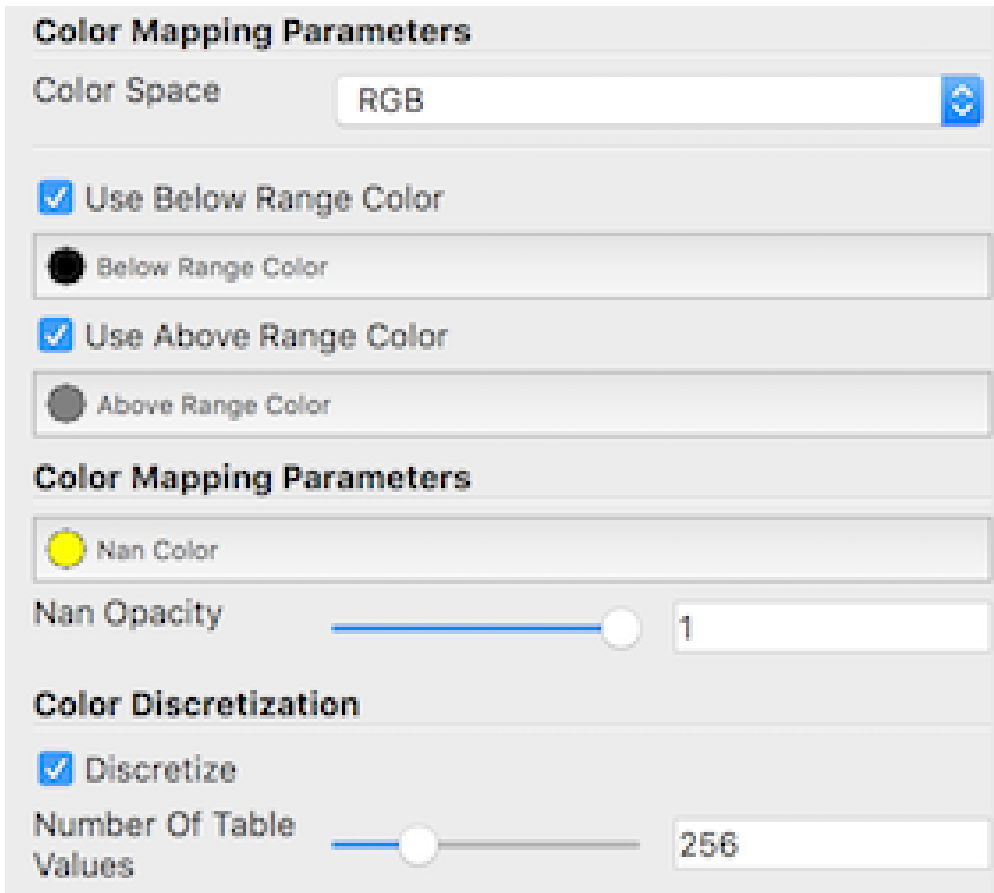


Fig. 2.7: Color Mapping Parameters, including advanced properties. Advanced properties are enabled by clicking the gear icon at the top right of the `Color Map Editor` (not shown.)

```

# Rescale transfer functions to a specific range
>>> colorMap.RescaleTransferFunction(1.0, 19.9495)
>>> opacityMap.RescaleTransferFunction(1.0, 19.9495)

# Invert the color map.
>>> colorMap.InvertTransferFunction()

# Map color map to log-scale preserving relative positions for
# control points
>>> colorMap.MapControlPointsToLogSpace()
>>> colorMap.UseLogScale = 1

# Return back to linear space.
>>> colorMap.MapControlPointsToLinearSpace()
>>> colorMap.UseLogScale = 0

# Change using of opacity mapping for surfaces
>>> colorMap.EnableOpacityMapping = 1

# Explicitly specify color map control points
# The value is a flattened list of tuples
# (data-value, red, green, blue). The color components
# must be in the range [0.0, 1.0]
>>> colorMap.RGBPoints = [1.0, 0.705, 0.015, 0.149,
                         5.0, 0.865, 0.865, 0.865,
                         10.0, 0.627, 0.749, 1.0,
                         19.9495, 0.231373, 0.298039, 0.752941]

# Similarly, for opacity map. The value here is
# a flattened list of (data-value, opacity, mid-point, sharpness)
>>> opacity.Points = [1.0, 0.0, 0.5, 0.0,
                      9.0, 0.404, 0.5, 0.0,
                      19.9495, 1.0, 0.5, 0.0]

# Note, in both these cases the controls points are assumed to be sorted
# based on the data values. Also, not setting the first and last
# control point to have same data value can have unexpected artifacts
# in the 'Color Map Editor' panel.

```

Oftentimes, you want to rescale the color and opacity maps to fit the current data ranges. You can do this as follows:

```

>>> source = GetActiveSource()

# Update the pipeline, if it hasn't been updated already.
>>> source.UpdatePipeline()

# First, locate the display properties for the source of interest.
>>> display = GetDisplayProperties()

# Reset the color and opacity maps currently used by 'display' to
# use the range for the array 'display' is using for color mapping.
# This requires that the 'display' has been set to use scalar coloring
# using an array that is available in the data generated. If not, you will

```

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```
# get errors.
>>> display.RescaleTransferFunctionToDataRange()
```

2.2.4 Color legend

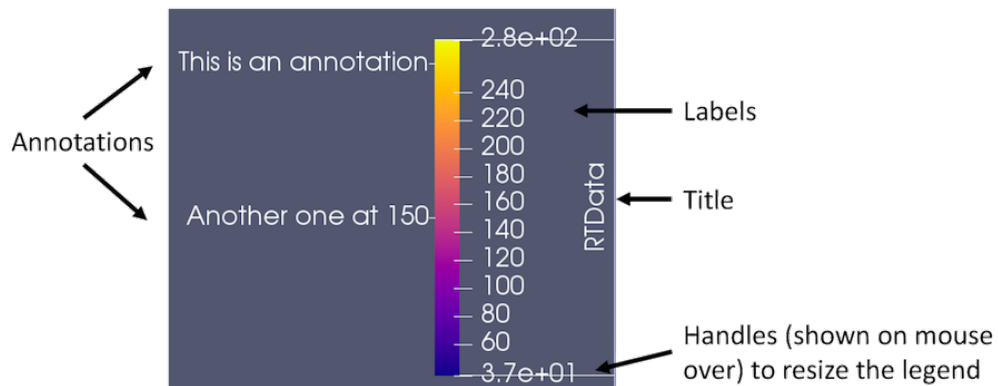


Fig. 2.8: Color legend in **ParaView**.

The color legend, also known as scalar bar or color bar, is designed to provide the user information about which color corresponds to what data value in the rendered view. You can toggle the visibility of the color legend corresponding to the transfer function being shown/edit in the **Color Map Editor** by using the  button near the top of the panel. This button affects the visibility of the legend in the active view.

Fig. 2.8 shows the various components of the color legend. By default, the title is typically the name of the array (and component number or magnitude for non-scalar data arrays) being mapped. Automatically generated labels appear on one side of the color legend, while on the other side are annotations, optionally including start and end annotations depicting the minimum and maximum of the color legend range.

The color legend can be manipulated with the mouse. You can click and drag the legend to place it at any position in the view. Additionally, you can change the length of the legend by clicking and dragging the end-markers shown when you hover the mouse pointer over the legend.

Color legend parameters

You can edit the color legend parameters by clicking on the  button on the **Color Map Editor** panel. This will pop up the **Edit Color Legend Properties** dialog that shows the available parameters. Any changes made will affect only the particular color legend in the active view.

The first few options in the **Edit Color Legend Properties** control the orientation and location of the color legend in the render view. **Auto Orient** turns on automatic determination of the color legend's orientation. The color legend will change orientation to horizontal when it is dragged to the bottom or the top of the render view, and it will change to vertical when it is dragged to the left or right side. When disabled, you can choose the orientation you want the color legend to have by choosing an option in the **Orientation** combo-box. The **Window Location** option controls the location of the color legend in the window; if the value **AnyLocation** is selected, then the color legend will not be forced into any particular position. The color legend can be positioned by clicking and dragging it with the mouse in the render view, or the **Position** property can be specified explicitly with fractional coordinates that range from [0, 1] and represent the fraction of the window width (or height) where the color legend's bottom left corner should be placed.

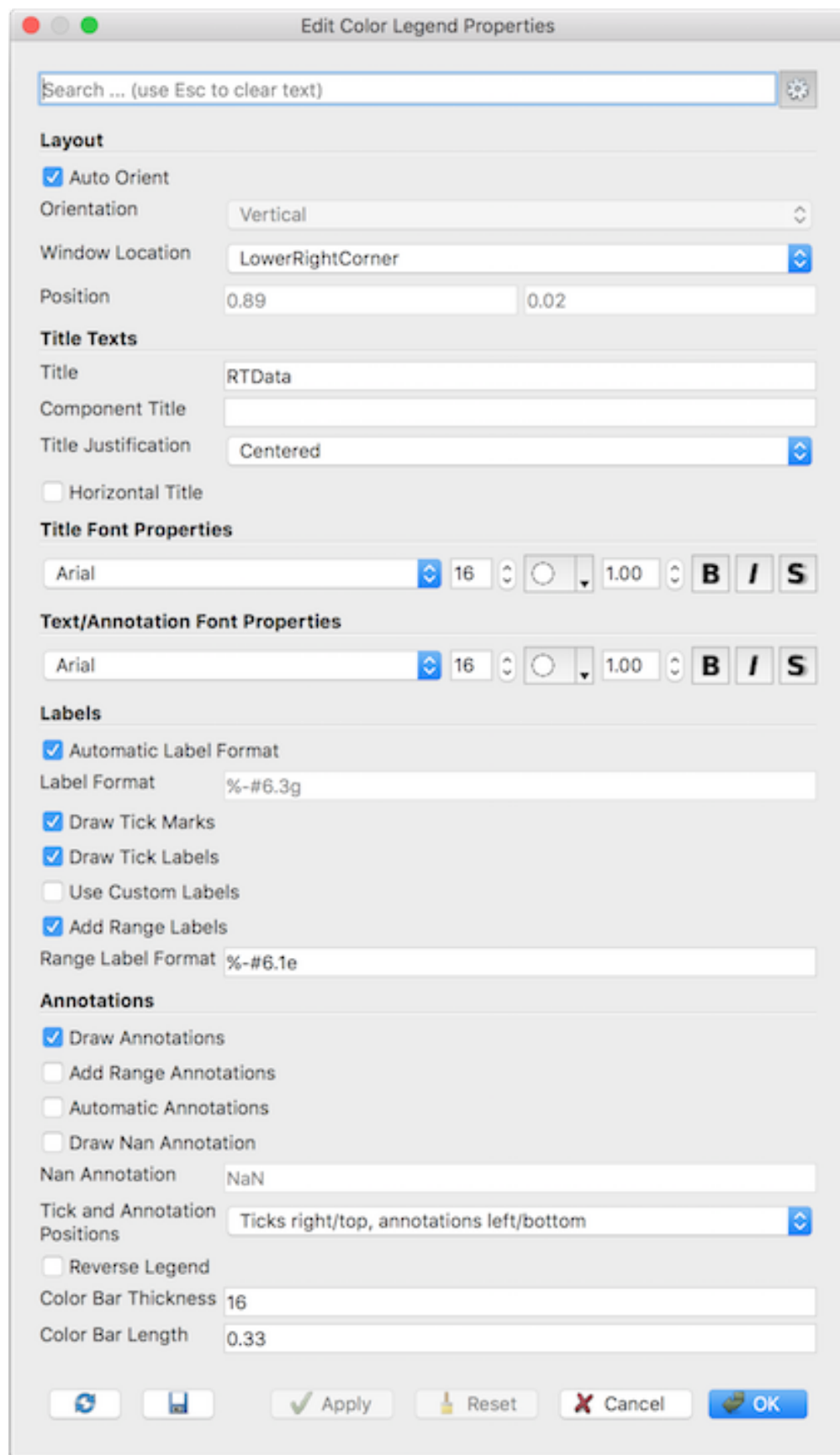


Fig. 2.9: Edit Color Legend Parameters dialog in paraview.
2.2. Color maps and transfer functions

Note that if the color legend is placed interactively with the mouse, the **Window Location** option will automatically change to **AnyLocation**.

Besides the obvious changing of title text and font properties for the title, labels, and annotations, there are some other parameters that control the appearance of the color legend.

By default, the title is rotated 90 degrees counter-clockwise when the legend is oriented vertically to better align with the legend. Checking the **Horizontal Title** box forces the title of the color legend to be horizontal regardless of color legend orientation.

Draw Annotations determines whether the annotations (including the start and end annotations) are to be drawn at all.

When checked, **Draw Nan Annotation** results in the color legend showing the NaN color set in the **Color Map Editor** panel in a separate color box right beside the color legend. The annotation text shown for that box can be modified by changing the **Nan Annotation** text.

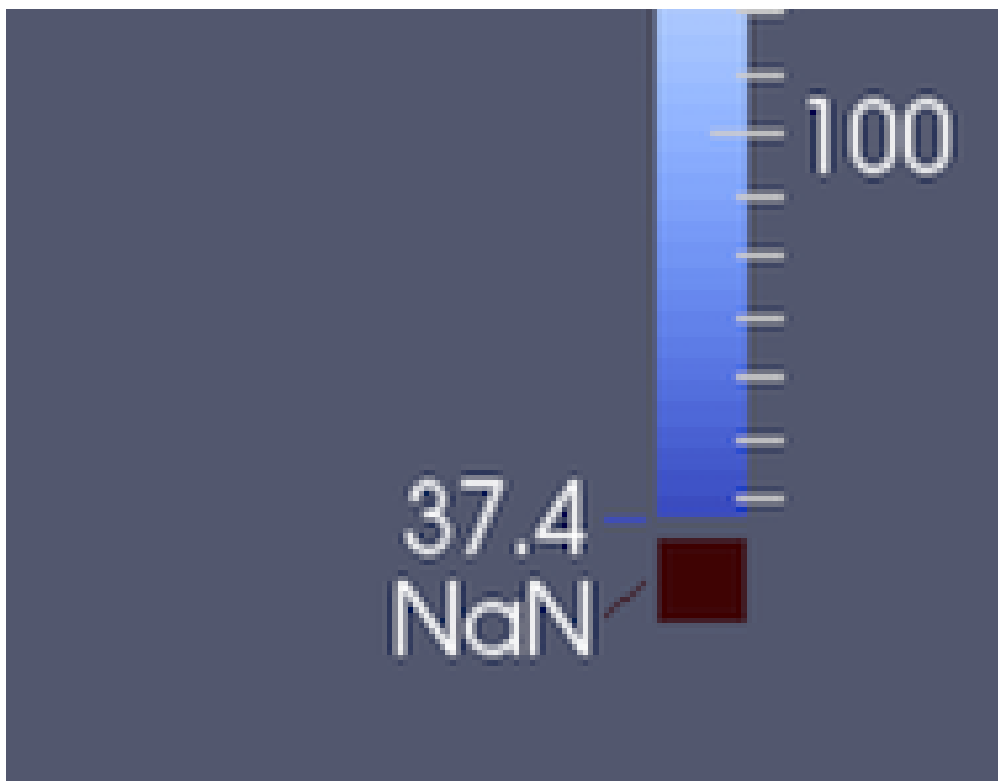


Fig. 2.10: Color legend showing NaN annotation

If **Automatic Label Format** is checked, **ParaView** will try to pick an optimal representation for numerical values based on the value and available screen space. By unchecking it, you can explicitly specify the printf-style format to use for numeric values. To explicitly label values of interest, enable the **Use Custom Labels** option. You can specify exactly the labeled values you wish to display in the table that appears when this option is chosen. **Color Bar Thickness** is used to control the thickness of the legend. It is defined in terms of points just like how font sizes are specified. Use **Color Bar Length** to explicitly set the length of the color bar. This property is defined as a fraction in the range [0, 1] of the window width (when the color legend is oriented horizontally) or height (when oriented vertically).

Color legend in ppython

To show the color legend or scalar bar for the transfer function used for scalar mapping a source in a view, you can use API on its display properties:

```
>>> source = ...
>>> display = GetDisplayProperties(source, view)

# to show the color legend
>>> display.SetScalarBarVisibility(view, True)

# to hide the same
>>> display.SetScalarBarVisibility(view, False)
```

To change the color legend properties as in Section 2.2.4, you need to first get access to the color legend object for the color transfer function in a particular view. These are analogous to display properties for a source in a view.

```
>>> colorMap = GetColorTransferFunction('Temp')

# get the scalar bar in a view (akin to GetDisplayProperties)
>>> scalarBar = GetScalarBar(colorMap, view)

# Now, you can change properties on the scalar bar object.
>>> scalarBar.TitleFontSize = 8
>>> scalarBar.DrawNaNAnnotation = 1
```

2.2.5 Annotations

Simply put, annotations allow users to put custom text at particular data values in the color legend. The min and max data mapped value annotations are automatically added. To add any other custom annotations, you can use the **Color Map Editor**.

Since the list of annotations is an advanced property, you need to either toggle the visibility of advanced properties using the icon near the top of the panel or type **annotations** in the search box. That will show the **Annotations** widget, which is basically a list widget where users can enter key-value pairs, rather than value-annotation pairs, as shown in Fig. 2.11.

You can use the buttons of the right on the widget to add/remove entries. Enter the data value to annotate under the **Value** column and then enter the text to display at that value under the **Annotation** column.

You can use the Tab (tab) key to edit the next entry. Hitting Tab after editing the last entry in the table will automatically result in adding a new row, thus, making it easier to add bunch of annotations without having to click any buttons.

Some annotation texts may not show up on the legend. There are two possible reasons an annotation may not be shown. First, the value added is outside the mapped range of the color transfer function. Second, **Draw Annotations** is unchecked in the **Color Legend Parameters** dialog .

The and buttons can be used to fill the annotations widget with unique discrete values from a data array, if possible. Based on the number of distinct values present in the data array, this may not yield any result (Instead, a warning message will be shown). The data array values come either from the selected source object if you use the button or it comes from the visible pipeline objects if you use the button.

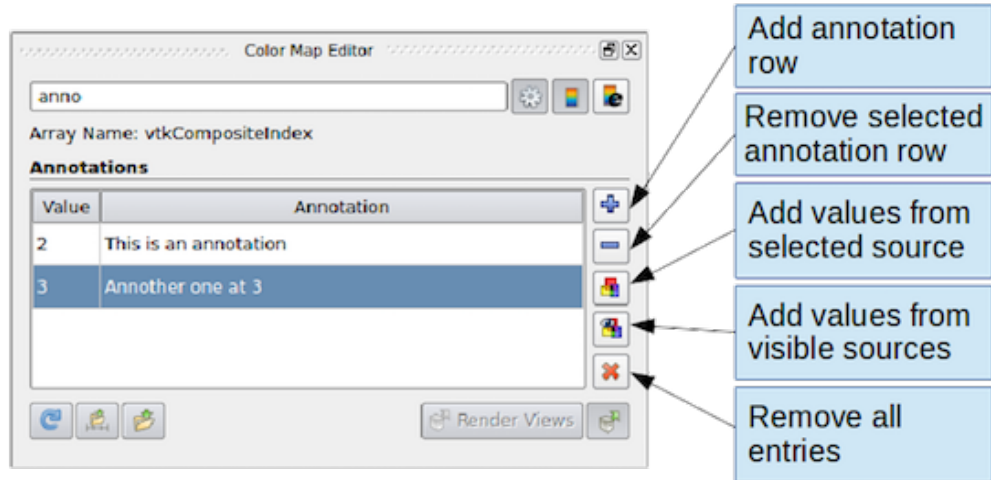


Fig. 2.11: Widget to add/edit annotations on the Color Map Editor panel

Annotations in pypython

Annotations is a property on the color map object. You simply get access to the color map of interest and then change the Annotations property.

```
>>> colorMap = GetColorTransferFunction('Temp')

# Annotations are specified as a flattened list of tuples
# (data-value, annotation-text)
>>> colorMap.Annotations = ['1', 'Slow',
                           '10', 'Fast']
```

2.2.6 Categorical colors

A picture is worth a thousand words, they say, so let's just let the picture do the talking. Categorical color maps allow you to render visualizations as shown in Fig. 2.12.

When one thinks of scalar coloring, one is typically talking of mapping numerical values to colors. However, in some cases, the numerical values are not really numbers, but enumerations such as element types and gear types (as in Fig. 2.12) or generally, speaking, categories. The traditional approach of using an interpolated color map specifying the range of values with which to color doesn't really work here. While users could always play tricks with the number of discrete steps, multiple control points, and annotations, it is tedious and cumbersome.

Categorical color maps provide an elegant solution for such cases. Instead of a continuous color transfer function, the user specifies a set of discrete values and colors to use for those values. For any element where the data value matches the values in the lookup table exactly, paraview renders the specified color; otherwise, the NaN color is used.

The color legend or scalar bar also switches to a new mode where it renders swatches with annotations, rather than a single bar. You can add custom annotations for each value in the lookup table.

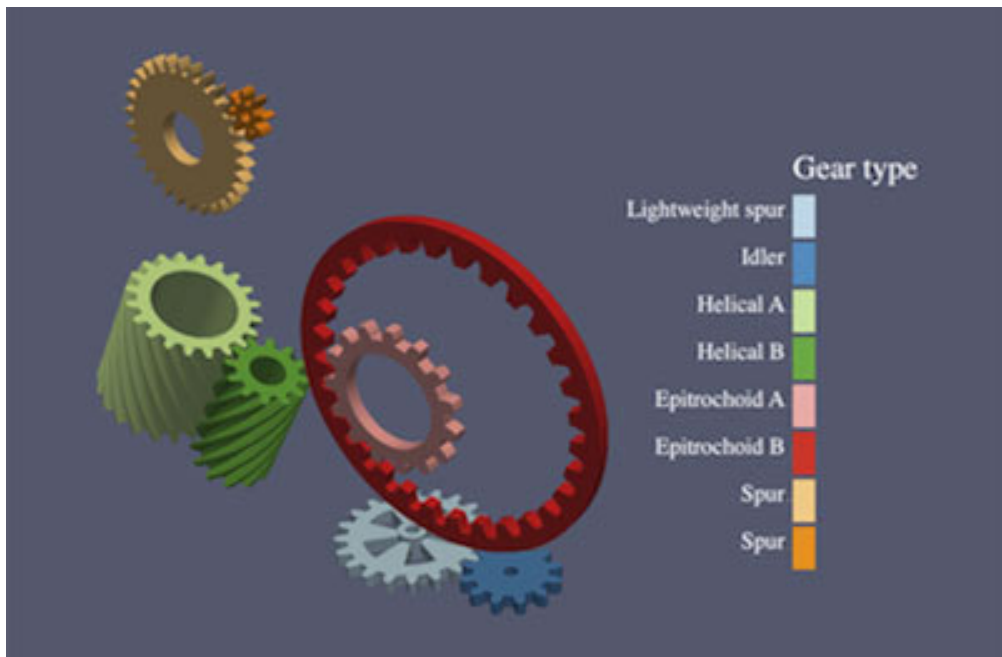


Fig. 2.12: Visualization using a categorical color map for discrete coloring

Categorical Color: User Interface

To tell paraview that the data array is to be treated as categories for coloring, check the `Interpret Values As Categories` checkbox in the `Color Map Editor` panel. As soon as that's checked, the panel switches to categorical mode: The `Mapping Data` group is hidden, and the `Annotations` group becomes a non-advanced group, i.e., the annotations widget is visible even if the panel is not showing advanced properties, as is shown in Fig. 2.13.

The annotations widget will still show any annotations that may have been added earlier, or it may be empty if none were added. You can add annotations for data values as was the case before using the buttons on the side of the widget. This time, however, each annotation entry also has a column for color. If color has not been specified, a question mark icon will show up; otherwise, a color swatch will be shown. You can double click the color swatch or the question mark icon to specify the color to use for that entry. Alternatively, you can choose from a preset collection of categorical color maps by clicking the  button.

As before, you can use `Tab` key to edit and add multiple values. Hence, you can first add all the values of interest in one pass and then pick a preset color map to set colors for the values added. If the preset has fewer colors than the annotated values, then the user may have to manually set the colors for those extra annotations.

Common Errors

Categorical color maps are designed for data arrays with enumerations, which are typically integer arrays. However, they can be used for arrays with floating point numbers as well. With floating point numbers, the value specified for annotation may not match the value in the dataset exactly, even when the user expects it to match. In that case, the `Nan` color will be used.

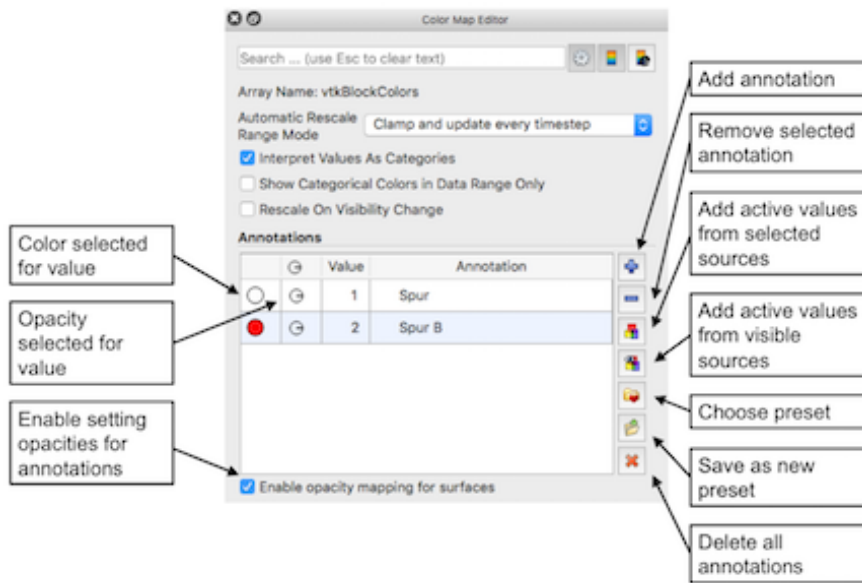


Fig. 2.13: Default Color Map Editor when Interpret Values As Categories is checked.

Categorical colors in pvython

```
>>> categoricalColorMap = GetColorTransferFunction('Modes')
>>> categoricalColorMap.InterpretValuesAsCategories = 1

# specify the labels for the categories. This is similar to how
# other annotations are specified.
>>> categoricalColorMap.Annotations = ['0', 'Alpha', '1', 'Beta']

# now, set the colors for each category. These are an ordered list
# of flattened tuples (red, green, blue). The first color gets used for
# the first annotation, second for second, and so on
>>> categoricalColorMap.IndexedColors = [0.0, 0.0, 0.0,
                                         0.89, 0.10, 0.10]
```

2.3 Comparative visualization

Comparative visualization in **ParaView** refers to the ability to create side-by-side visualizations for comparing with one another. In its most basic form, you can indeed use **ParaView**'s ability to show multiple views side by side to set up simultaneous visualizations. But, that can get cumbersome too quickly. Let's take a look at a simple example: Let's say you want to do a parameter study where you want to compare isosurfaces generated by a set of isovalue in a dataset. To set up such a visualization, you'll need to first create as many **Render View**s as isovalue. Then, create just as many **Contour** filters, setting each one up with a right isovalue for the contour to generate and display the result in one of the views. As the number of isovalue increases, you can see how this can get tedious. It is highly error prone, as you need to keep track of which view shows which isovalue. Now, imagine after having set up the entire visualization that you need to change the **Representation** type for all the of the isosurfaces to **Wireframe**!

Comparative Views were designed to handle such use-cases. Instead of creating separate views, you create a single view **Render View (Comparative)**. The view itself comprises of a configurable $m \times n$ **Render View**s. Any data

that you show in this view gets shown in all the internal views simultaneously. Any display property changes, such as scalar coloring and representation type are also maintained consistently between these internal views. The interactions in the internal views are linked, so when you interact with one, all other views update as well. While this is all nice and well, the real power of Comparative View s becomes apparent when you set up a parameter to vary across the views. This parameter can be any of the properties on the pipeline modules such as filter properties and opacity, or it could be the data time. Each of these internal views will now render the result obtained by setting the parameter as per your selection. Going back to our original example, we will create a single Contour filter that we show in Render View (Comparative) with as many internal views as the isovalue to compare. Next, we will set up a parameter study for varying the Isosurfaces property on the Contour filter, and, viola! The view will generate the comparative visualization for us!

In this chapter, we look at how to configure this view and how to set these parameters to compare. We limit our discussion to Render View (Comparative) . However, the same principles are applicable to other comparative views, including Bar Chart View (Comparative) and Line Chart View (Comparative) .

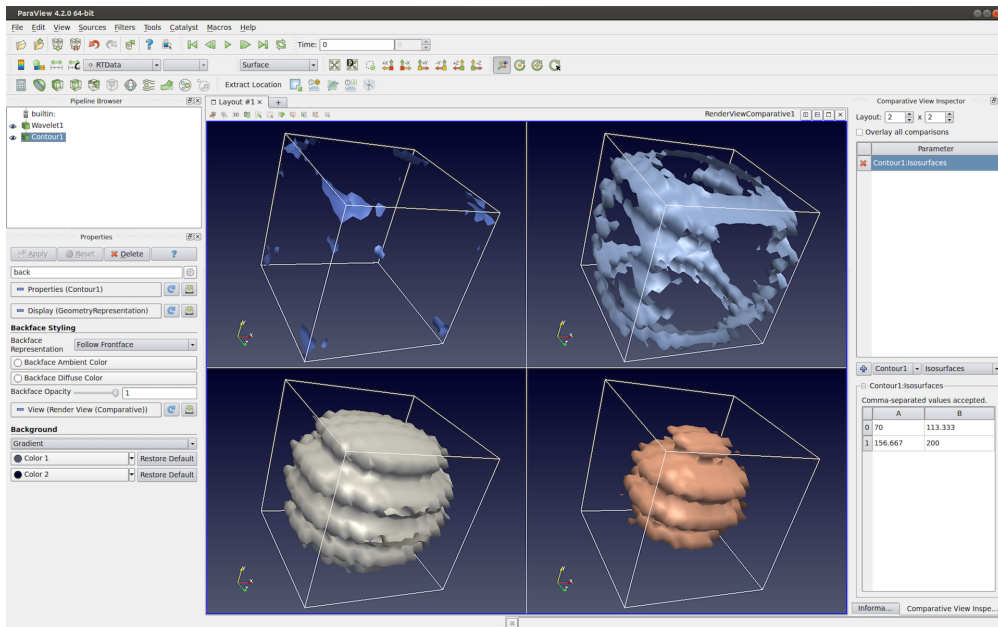


Fig. 2.14: Render View (Comparative) in paraview showing a parameter study. In this case, we are comparing the visualization generated using different isovalue s for the Contour filter. The Comparative View Inspector dockpanel (on the right) is used to configure the parameter study.

2.3.1 Setting up a comparative view

To create Render View (Comparative) in paraview, split or close the active view and select Render View (Comparative) from the new view creation widget. paraview will immediately show four Render View s laid out in a 2×2 grid. While you cannot resize these internal views, notice that you can still split the view frame and create other views if needed.

The Properties panel will show properties similar to those available on the Render View under the View properties section. If you change any of these properties, they will affect all these internal views, e.g., setting the Background color to Gradient will make all the views show a gradient background.

To configure the comparative view itself, you must to use the Comparative View Inspector (Fig. 2.15) accessible from the View menu. The Comparative View Inspector is a dockable panel that is enabled when the active view is a comparative view.

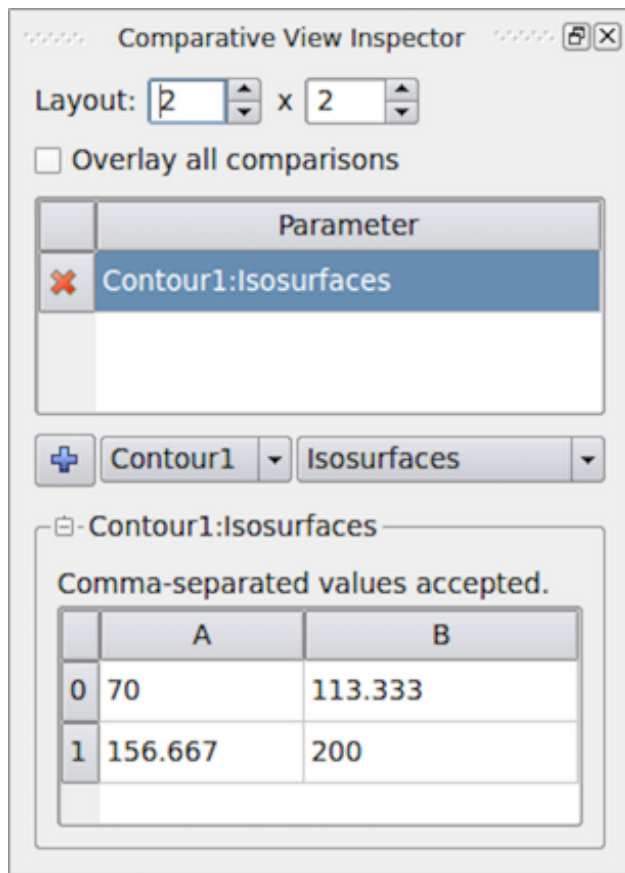


Fig. 2.15: Comparative View Inspector in paraview used to configure the active comparative view.

To change how many internal views are used in the active comparative view and how they are laid out, use the **Layout**. The first value is the number of views in the horizontal direction and the second is the count in the vertical direction.

Besides doing a parameter study in side-by-side views, you can also show all the results in a single view. For that, simply check the **Overlay all comparisons** checkbox. When checked, instead of getting a grid of $m \times n$ views, you will only see one view with all visible data displayed $m \times n$ times.

To show data in this view is the same as any other view: Make this view active and use the **Pipeline Browser** to toggle the eyeball icons to show data produced by the selected pipeline module(s). Showing any dataset in this view will result in the data being shown in all the internal views simultaneously. As is true with **View properties**, with **Display properties**, changing **Coloring**, **Styling**, or any other properties will reflect in all internal views.

Since the cameras among the internal views are automatically linked, if you interact with one of the views, all views will also update simultaneously when you release the mouse button.

2.3.2 Setting up a parameter study

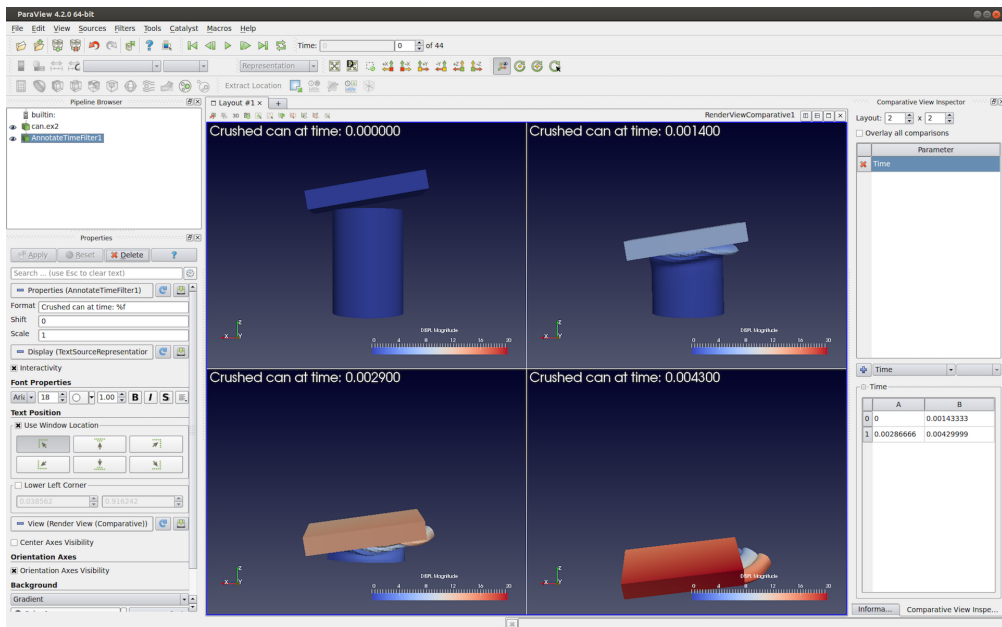


Fig. 2.16: Render View (Comparative) with **Annotate Time** filter showing the time for each of the views. In this case, the parameter study is varying **Time** across the views.

To understand how to setup a parameter study, let's go back to our original example. Our visualization pipeline is simply **Wavelet** → **Contour** (assuming default property values), and we are showing the result from the **Contour** filter in the 2×2 Render View (Comparative) with **Overlay all comparisons** unchecked.

Now, we want to vary the **Isosurfaces** property value for the visualization in each of the internal views. That's our parameter to study. Since that's a property on the **Contour** filter, we select the **Contour** filter instance. Then, its property **Isosurfaces** is in the parameter selection combo-boxes. To add the parameter to the study, you must hit the button.

The parameter **Contour1: Isosurfaces** will then show up in the Parameter list above the combo-boxes. You can delete this parameter by using the button next to the parameter name.

Did you know?

Notice how this mechanism for setting up a parameter study is similar to how animations are set up. In fact, under the cover, the two mechanisms are not that different, and they share a lot of implementation code.

Simultaneously, the table widget below the combo-boxes is populated with a default set of values for the `Isosurfaces`. This is where you specify the parameter values for each of the views. The location of the views matches the location in the table. Hence, $0 - A$ is the top-left view, $0 - B$ is the second view from the left in the topmost row, and so on. The parameter value at a specific location in the table is the value used to generate the visualization in the corresponding view.

To change the parameter (in our case `Isosurfaces`) value, you can double click on the cell in the table and change it one at a time. Also, to make it easier to fill the cells with a range of values, you can click and drag over multiple cells. When you release the mouse, a dialog will prompt you to enter the data value range ([Fig. 2.17](#)). In case you selected a combination of rows and columns, the dialog will also let you select in which direction the parameter is varied using the range specified. You can choose to vary the parameter horizontally first, vertically first, or only along one of the directions while keeping the other constant. This is useful when doing a study with multiple parameters.

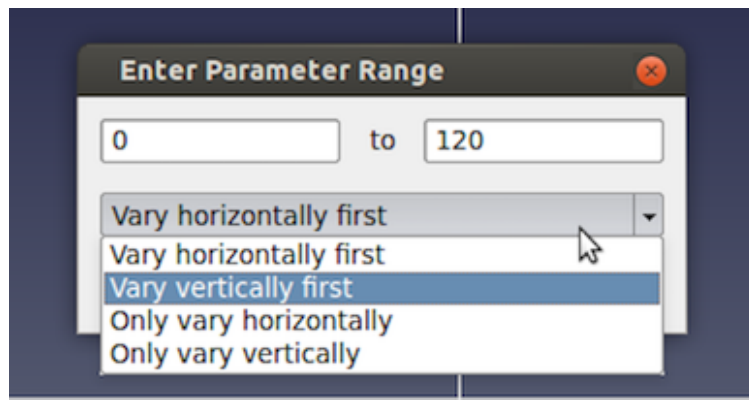


Fig. 2.17: Dialog used to select a range of parameter values and control how to vary them in paraview.

As soon as you change the parameter values, the view will update to reflect the change.

2.3.3 Adding annotations

You add annotations like color legends, text, and cube-axes exactly as you would with a regular `Render View`. As with other properties, annotations will show up in all of the internal views.

Did you know?

You can use the `Annotate Time` source or filter to show the data time or the view time in each of the internal views. If `Time` is one of the parameters in your study, the text will reflect the time values for each of the individual views, as expected ([Fig. 2.16](#))!

2.4 Programmable Filter

A pipeline module in **ParaView** does one of two things: it either generates data or processes input data. To generate data, the module may use a mathematical model e.g. *Sources > Sphere* or read a file from disk. Processing data entails transforming input data by applying defined operations to generate a new output. **ParaView** provides a large set of readers, data sources and data filters that cover the needs of many users. For the cases where the available collection does not satisfy your needs, **ParaView** provides a mechanism to add new modules via plugins. Conventional plugins, however, are intended for hardcore developers. They are written in C++, using the data processing APIs provided by **ParaView** and VTK. The complexity of building and packaging C++ plugins that work on all distributed versions of **ParaView** can be daunting and thus a huge barrier for even advanced **ParaView** users. Python-based programmable filters, sources and annotations provide an easy alternative to this. New modules can be written as Python scripts that are executed by **ParaView** to generate, process and/or display data, just like conventional C++ modules. Since the scripts are standard Python scripts, you have access to Python packages such as NumPy that provide several numeric operations useful for data transformation.

In this chapter, we will explore how to use Python to add new data processing modules to ParaView through examples. For additional explanation of the data processing API, see [Section 2.5](#).

Common Errors

In this guide so far, we have been looking at examples of Python scripts for `pvpython`. These scripts are used to script the actions you would perform using the `paraview` UI. The scripts you would write for **Programmable Source**, **Programmable Filter**, and **Programmable Annotation** are entirely different. The data processing API executes within the data processing pipeline and, thus, has access to the data being processed. In client-server mode, this means that such scripts are indeed executed on the server side, potentially in parallel, across several MPI ranks. Therefore, attempting to import the `paraview.simple` Python module in the **Programmable Source** script, for example, is not supported and will have unexpected consequences.

2.4.1 Understanding the programmable modules

With programmable modules, you are writing custom code for filters and sources. You are expected to understand the basics of a VTK (and **ParaView**) data processing pipeline, including the various stages of the pipeline execution as well as the data model. Refer to [Section 1.3.1](#) for an overview of the VTK data model. While a detailed discussion of the VTK pipeline execution model is beyond the scope of this book, the fundamentals covered in this section, along with the examples in the rest of this chapter, should help you get started and write useful modules. For a primer on the details of the VTK pipeline execution stages, see [\[BerkGeveci\]](#).

To create the programmable source, filter, or annotation in `paraview`, you use the *Sources > Programmable Source*, *Filters > Programmable Filter* or *Sources > Programmable Annotation* menus, respectively. Since the **Programmable Filter** and **Programmable Annotation** are filters, like other filters, they get connected to the currently active source(s), i.e., the currently active source(s) become the input to this new filter. **Programmable Source**, on the other hand, does not have any inputs.

One of the first things that you specify after creating a programmable filter or source is the **Output Data Set Type**. This option lets you select the type of dataset this module will produce. The options provided include several of the data types discussed in [Section 1.3.1](#). Additionally, for the **Programmable Filter**, you can select **Same as Input** to indicate that the filter preserves the input dataset type.

Next is the primary part: the **Script**. This is where you enter the Python script to generate or process from the inputs the dataset that the module will produce. As with any Python script, you can import other Python packages and modules in this script. Just be aware that when running in client-server mode, this script is going to be executed on the server side. Accordingly, any modules or packages you import must be available on the server side to avoid errors.

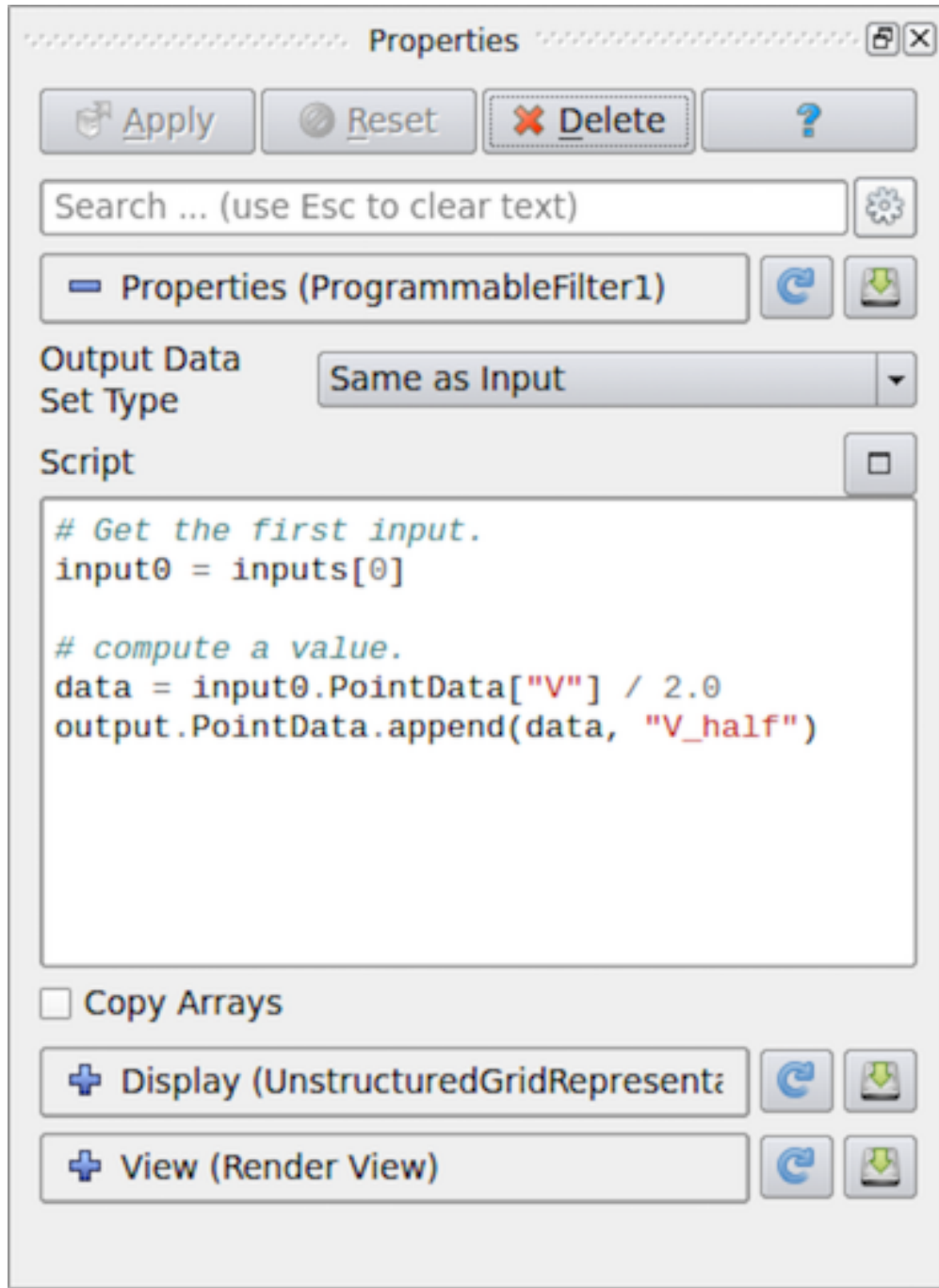


Fig. 2.18: Properties panel for Programmable Filter in paraview.

The script gets executed in what's called the *RequestData* pass of the pipeline execution. This is the pipeline pass in which an algorithm is expected to produce the output dataset.

There are several other passes in a pipeline's execution. The ones for which you can specify a Python script to execute in the programmable filter and source are:

- *RequestInformation*: In this pass, the algorithm is expected to provide the pipeline with any meta-data available about the data that will be produced by it. This includes things like number of timesteps in the dataset and their time values for temporal datasets or extents for structured datasets. This gets called before *RequestData* pass. In the *RequestData* pass, the pipeline could potentially qualify the request based on the meta-data provided in this pass. For example, if an algorithm announces that the temporal dataset has multiple timesteps, the pipeline could request that the algorithm produce data for one of those timesteps in *RequestData*.
- *RequestUpdateExtent*: In this pass, a filter gets the opportunity to qualify requests for execution passed on to the upstream pipeline. As an example, if an upstream reader announced in its *RequestInformation* script that it can produce several timesteps, in *RequestUpdateExtent*, this filter can make a request to the upstream reader for a specific timestep. This pass gets called after *RequestInformation*, but before *RequestData*. It's not very common to provide a script for this pass.

You can specify the script for the *RequestInformation* pass in `RequestInformation Script` and for the *RequestUpdateExtent* pass in `RequestUpdateExtent Script`. Since the *RequestUpdateExtent* pass does not make much sense for an algorithm that does not have any inputs, `RequestUpdateExtent Script` is not available on `Programmable Source`. `Programmable Annotation` only has a *RequestData* script as this is the only one that make sense in this context.

2.4.2 Recipes for Programmable Source

In this section, we look at several recipes for `Programmable Source`. A common use of `Programmable Source` is to prototype readers. If your reader library already provides a Python API, then you can easily import the appropriate Python package to read your dataset using `Programmable Source`.

Did you know?

Most of the examples in this chapter use a NumPy-centric API for accessing and creating data arrays. Additionally, you can use VTK's Python wrapped API for creating and accessing arrays. Given the omnipresence of NumPy, there is rarely any need for using VTK's API directly, however.

Reading a CSV file

For this example, we will read in a CSV file to produce a Table (Section 1.3.1) using `Programmable Source`. We will use NumPy to do the parsing of the CSV files and pass the arrays read in directly to the pipeline. Note that the `Output DataSet Type` must be set to `vtkTable`.

```
# Code for 'Script'

# We will use NumPy to read the csv file.
# Refer to NumPy documentation for genfromtxt() for details on
# customizing the CSV file parsing.

import numpy as np
# assuming data.csv is a CSV file with the 1st row being the names names for
# the columns
data = np.genfromtxt("data.csv", dtype=None, names=True, delimiter=',', autostrip=True)
```

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```

for name in data.dtype.names:
    array = data[name]

    # You can directly pass a NumPy array to the pipeline.
    # Since ParaView expects all arrays to be named, you
    # need to assign it a name in the 'append' call.
    output.RowData.append(array, name)

```

Reading a CSV file series

Building on the example from [Section 2.4.2](#), let's say we have a series of files that we want to read in as a temporal series. Recall from [Section 2.4.1](#) that meta-data about data to be produced, including timestep information, is announced in `RequestInformation` pass. Hence, for this example, we will need to specify the `RequestInformation Script` as well.

As was true earlier, `Output DataSet Type` must be set to `vtkTable`. Now, to announce the timesteps, we use the following as the `RequestInformation Script`.

```

# Code for 'RequestInformation Script'.
def setOutputTimesteps(algorithm, timesteps):
    "helper routine to set timestep information"
    executive = algorithm.GetExecutive()
    outInfo = executive.GetOutputInformation(0)

    outInfo.Remove(executive.TIME_STEPS())
    for timestep in timesteps:
        outInfo.Append(executive.TIME_STEPS(), timestep)

    outInfo.Remove(executive.TIME_RANGE())
    outInfo.Append(executive.TIME_RANGE(), timesteps[0])
    outInfo.Append(executive.TIME_RANGE(), timesteps[-1])

# As an example, let's say we have 4 files in the file series that we
# want to say are producing time 0, 10, 20, and 30.
setOutputTimesteps(self, (0, 10, 20, 30))

```

The Script is similar to earlier, except that we will read a specific CSV file based on which timestep was requested.

```

# Code for 'Script'
def GetUpdateTimestep(algorithm):
    """Returns the requested time value, or None if not present"""
    executive = algorithm.GetExecutive()
    outInfo = executive.GetOutputInformation(0)
    return outInfo.Get(executive.UPDATE_TIME_STEP()) \
        if outInfo.Has(executive.UPDATE_TIME_STEP()) else None

# This is the requested time-step. This may not be exactly equal to the
# timesteps published in RequestInformation(). Your code must handle that
# correctly.
req_time = GetUpdateTimestep(self)

# Now, use req_time to determine which CSV file to read and read it as before.

```

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```
# Remember req_time need not match the time values put out in
# 'RequestInformation Script'. Your code need to pick an appropriate file to
# read, irrespective.

...
# TODO: Generate the data as you want.

# Now mark the timestep produced.
output.GetInformation().Set(output.DATA_TIME_STEP(), req_time)
```

Reading a CSV file with particles

This is similar to Section 2.4.2. Now, however, let's say the CSV has three columns named X, Y and Z that we want to treat as point coordinates and produce a vtkPolyData with points instead of a vtkTable . For that, we first ensure that Output DataSet Type is set to vtkPolyData . Next, we use the following Script :

```
# Code for 'Script'

from vtk.numpy_interface import algorithms as algs
from vtk.numpy_interface import dataset_adapter as dsa
import numpy as np

# assuming data.csv is a CSV file with the 1st row being the names names for
# the columns
data = np.genfromtxt("/tmp/points.csv", dtype=None, names=True, delimiter=',', ,
                     autostrip=True)

# convert the 3 arrays into a single 3 component array for
# use as the coordinates for the points.
coordinates = algs.make_vector(data["X"], data["Y"], data["Z"])

# create a vtkPoints container to store all the
# point coordinates.
pts = vtk.vtkPoints()

# numpyToVtkDataArray is needed to called directly to convert the NumPy
# to a vtkDataArray which vtkPoints::SetData() expects.
pts.SetData(dsa.numpyToVtkDataArray(coordinates, "Points"))

# set the pts on the output.
output.SetPoints(pts)

# next, we define the cells i.e. the connectivity for this mesh.
# here, we are creating merely a point could, so we'll add
# that as a single poly vertex cell.
numPts = pts.GetNumberOfPoints()
# ptIds is the list of point ids in this cell
# (which is all the points)
ptIds = vtk.vtkIdList()
ptIds.SetNumberOfIds(numPts)
for a in xrange(numPts):
```

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```

ptIds.SetId(a, a)

# Allocate space for 1 cell.
output.Allocate(1)
output.InsertNextCell(vtk.VTK_POLY_VERTEX, ptIds)

# We can also pass all the array read from the CSV
# as point data arrays.
for name in data.dtype.names:
    array = data[name]
    output.PointData.append(array, name)

```

The thing to note is that this time, we need to define the geometry and topology for the output dataset. Each data type has different requirements on how these are specified. For example, for unstructured datasets like vtkUnstructuredGrid and vtkPolyData, we need to explicitly specify the geometry and all the connectivity information. For vtkImageData, the geometry is defined using origin, spacing, and extents, and connectivity is implicit.

Reading binary 2D image

This recipe shows how to read raw binary data representing a 3D volume. Since raw binary files don't encode information about the volume extents and data type, we will assume that the extents and data type are known and fixed.

For producing image volumes, you need to provide the information about the structured extents in *RequestInformation*. Ensure that the Output Data Set Type is set to `vtkImageData`.

```

# Code for 'RequestInformation Script'.
executive = self.GetExecutive()
outInfo = executive.GetOutputInformation(0)
# we assume the dimensions are (48, 62, 42).
outInfo.Set(executive.WHOLE_EXTENT(), 0, 47, 0, 61, 0, 41)
outInfo.Set(vtk.vtkDataObject.SPACING(), 1, 1, 1)

```

The Script to read the data can be written as follows.

```

# Code for 'Script'
import numpy as np

# read raw binary data.
# ensure 'dtype' is set properly.
data = np.fromfile("HeadMRVolume.raw", dtype=np.uint8)

dims = [48, 62, 42]
assert data.shape[0] == dims[0]*dims[1]*dims[2], "dimension mismatch"

output.SetExtent(0, dims[0]-1, 0, dims[1]-1, 0, dims[2]-1)
output.PointData.append(data, "scalars")
output.PointData.SetActiveScalars("scalars")

```

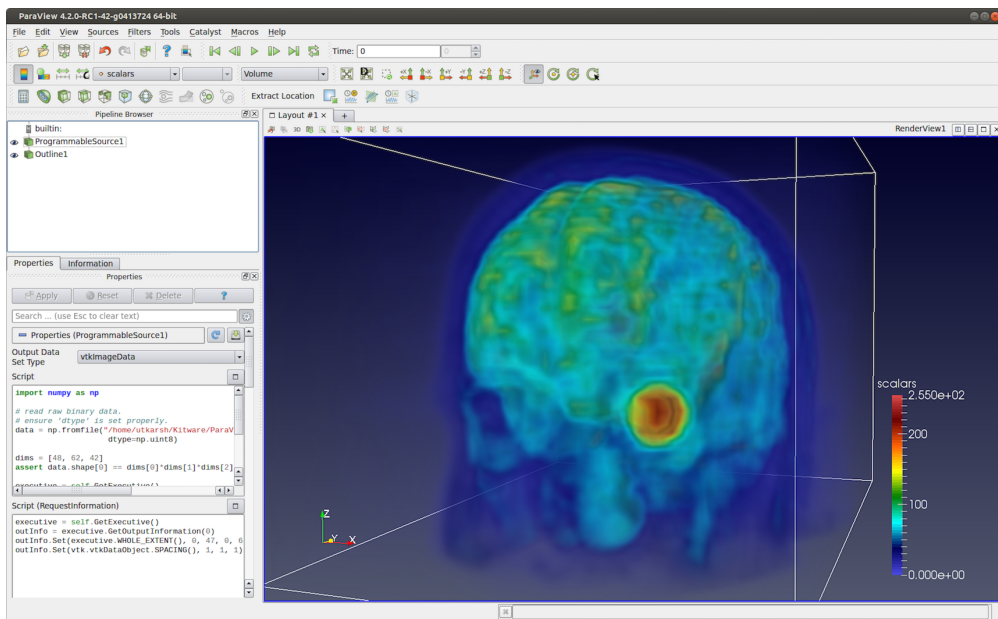


Fig. 2.19: Programmable Source used to read *HeadMRVolume.raw* file available in the VTK data repository.

Helix source

Here is another polydata source example. This time, we generate the data programmatically.

```
# Code for 'Script'

#This script generates a helix curve.
#This is intended as the script of a 'Programmable Source'
import math
import numpy as np
from vtk.numpy_interface import algorithms as algs
from vtk.numpy_interface import dataset_adapter as dsa

numPts = 80 # Points along Helix
length = 8.0 # Length of Helix
rounds = 3.0 # Number of times around

# Compute the point coordinates for the helix.
index = np.arange(0, numPts, dtype=np.int32)
scalars = index * rounds * 2 * math.pi / numPts
x = index * length / numPts;
y = np.sin(scalars)
z = np.cos(scalars)

# Create a (x,y,z) coordinates array and associate that with
# points to pass to the output dataset.
coordinates = algs.make_vector(x, y, z)
pts = vtk.vtkPoints()
pts.SetData(dsa.numpyToVtkDataArray(coordinates, 'Points'))
output.SetPoints(pts)
```

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```

# Add scalars to the output point data.
output.PointData.append(index, 'Index')
output.PointData.append(scalars, 'Scalars')

# Next, we need to define the topology i.e.
# cell information. This helix will be a single
# polyline connecting all the points in order.
ptIds = vtk.vtkIdList()
ptIds.SetNumberOfIds(numPts)
for i in xrange(numPts):
    #Add the points to the line. The first value indicates
    #the order of the point on the line. The second value
    #is a reference to a point in a vtkPoints object. Depends
    #on the order that Points were added to vtkPoints object.
    #Note that this will not be associated with actual points
    #until it is added to a vtkPolyData object which holds a
    #vtkPoints object.
    ptIds.SetId(i, i)

# Allocate the number of 'cells' that will be added. We are just
# adding one vtkPolyLine 'cell' to the vtkPolyData object.
output.Allocate(1, 1)

# Add the poly line 'cell' to the vtkPolyData object.
output.InsertNextCell(vtk.VTK_POLY_LINE, ptIds)

```

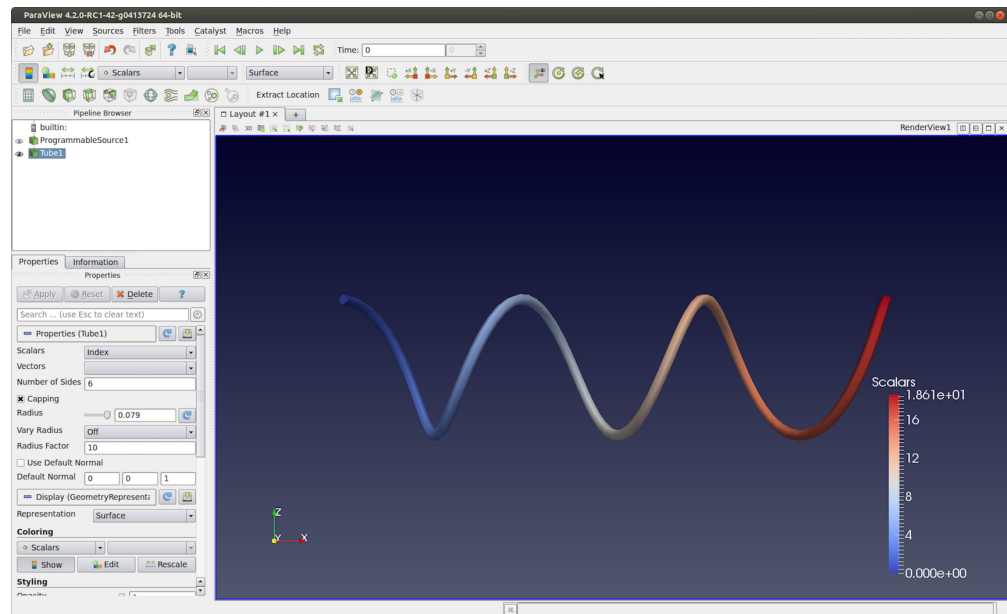


Fig. 2.20: Programmable Source output generated using the script in Section 2.4.2. This visualization uses Tube filter to make the output polyline more prominent.

2.4.3 Recipes for Programmable Filter

One of the differences between the Programmable Source and the Programmable Filter is that the latter expects at least 1 input. Of course, the code in the Programmable Filter is free to disregard the input entirely and work exactly as Programmable Source . Programmable Filter is designed to customize data transformations. For example, it is useful when you want to compute derived quantities using expressions not directly possible with Python Calculator and Calculator or when you want to use other Python packages or even VTK filters not exposed in ParaView for processing the inputs.

In this section, we look at various recipes for Programmable Filter s.

Adding a new point/cell data array based on an input array

Python Calculator provides an easy mechanism of computing derived variables. You can also use the Programmable Filter . Typically, for such cases, ensure that the Output DataSet Type is set to Same as Input .

```
# Code for 'Script'

# 'inputs' is set to an array with data objects produced by inputs to
# this filter.

# Get the first input.
input0 = inputs[0]

# compute a value.
dataArray = input0.PointData["V"] / 2.0

# To access cell data, you can use input0.CellData.

# 'output' is a variable set to the output dataset.
output.PointData.append(dataArray, "V_half")
```

The thing to note about this code is that it will work as expected even when the input dataset is a composite dataset such as a Multiblock dataset (Section 1.3.1). Refer to Section 2.5 for details on how this works. There are cases, however, when you may want to explicitly iterate over blocks in an input multiblock dataset. For that, you can use the following snippet.

```
input0 = inputs[0]
if input0.IsA("vtkCompositeDataSet"):
    # iterate over all non-empty blocks in the input
    # composite dataset, including multiblock and AMR datasets.
    for block in input0:
        processBlock(block)
else:
    processBlock(input0)
```

Computing tetrahedra volume

This recipe computes the volume for each tetrahedral cell in the input dataset. You can always simply use the Python Calculator to compute cell volume using the expression `volume(inputs[0])`. This recipe is provided to illustrate the API.

Ensure that the output type is set to Same as Input, and this filter assumes the input is an unstructured grid (Section 1.3.1).

```
# Code for 'Script'.

import numpy as np

# This filter computes the volume of the tetrahedra in an unstructured mesh.
# Note, this is just an illustration and not the most efficient way for
# computing cell volume. You should use 'Python Calculator' instead.
input0 = inputs[0]

numTets = input0.GetNumberOfCells()

volumeArray = np.empty(numTets, dtype=np.float64)
for i in xrange(numTets):
    cell = input0.GetCell(i)
    p1 = input0.GetPoint(cell.GetPointId(0))
    p2 = input0.GetPoint(cell.GetPointId(1))
    p3 = input0.GetPoint(cell.GetPointId(2))
    p4 = input0.GetPoint(cell.GetPointId(3))
    volumeArray[i] = vtk.vtkTetra.ComputeVolume(p1,p2,p3,p4)

output.CellData.append(volumeArray, "Volume")
```

Labeling common points between two datasets

In this example, the Programmable Filter takes two input datasets: A and B. It outputs dataset B with a new scalar array that labels the points in B that are also in A. You should select two datasets in the pipeline browser and then apply the programmable filter.

```
# Code for 'Script'

# Get the two inputs
A = inputs[0]
B = inputs[1]
# use len(inputs) to determine how many inputs are connected
# to this filter.

# We use numpy.in1d to test all which point coordinate components
# in B are present in A as well.
maskX = np.in1d(B.Points[:,0], A.Points[:,0])
maskY = np.in1d(B.Points[:,1], A.Points[:,1])
maskZ = np.in1d(B.Points[:,2], A.Points[:,2])

# Combining each component mask, we get the mask for point
# itself.
```

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```

mask = maskX & maskY & maskZ

# Now convert it to uint8, since bool arrays
# cannot be passed back to the VTK pipeline.
mask = np.asarray(mask, dtype=np.uint8)

# Initialize the output and add the labels array

# This ShallowCopy is needed since by default the output is
# initialized to be a shallow copy of the first input (inputs[0]),
# but we want it to be a description of the second input.
output.ShallowCopy(B.VTKObject)
output.PointData.append(mask, "labels")

```

Note in the script above the two inputs are defined in an `inputs` array. The order of elements in this array is determined by the order the data sources were selected in the Pipeline Browser . Hence, `inputs[0]` is the first data source selected and `inputs[1]` is the second.

2.4.4 Recipes for Programmable Annotation

The main difference between the programmable annotation and other programmable modules is that output is intended to be shown as a text representation. The output is expected to be a `vtkTable` containing a single one-component string array containing a single tuple. This tuple will be shown as a text representation, similar to the output of the Text source or the Python Annotation filter.

By default, the `Programmable Annotation` script already contains all that is needed to create this table, it just need to be filled up.

Similar to the `Programmable Filter`, the `Programmable Annotation` is designed to customize data display. For example, it is useful when you want to compute derived quantities using expressions not directly possible with Python Annotation and other annotation filters or when you want to use other Python packages or even VTK filters not exposed in `ParaView` for processing the inputs.

In this section, we look at various recipes for `Programmable Annotation`s.

Displaying the number of cells with a non-zero volume

`Python Annotation` provides easy mechanism to compute many values on datasets and arrays, but conditional operation like the one proposed here require the use of a `Programmable Annotation`.

```

# Code for 'Script'

# 'inputs' is set to an array with data objects produced by inputs to
# this filter.

# Get the first input.
input0 = inputs[0]

# compute the volume of each cell of the input
vols = volume(input0)

# the codepath for composite dataset and non composite dataset can't be shared

```

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```

# with this operation
if input0.IsA("vtkCompositeDataSet"):

    # create a running sum to iterate over blocks
    num = 0;

    # iterate over blocks
    for i in range(size(vols.Arrays)):
        # count the number of cells with a non-zero volume in this block
        # and add it to the running sum
        num += sum(val > 0 for val in vols.Arrays[i])
else:
    # non-composite case : just count the number of cells with a non-zero volume
    num = sum(val > 0 for val in vols)

# standard code to display the result
to = self.GetTableOutput()
arr = vtk.vtkStringArray()
arr.SetName("Text")
arr.SetNumberOfComponents(1)
arr.InsertNextValue(str(num))
to.AddColumn(arr)

```

Display system and date information

Python Annotation does not provide any mechanism to import Python modules while doing that in a Programmable Annotation is trivial.

```

# Code for 'Script'

# import needed python modules
from datetime import date
import platform

# construct the string to display
string = "Date: %s\n" % date.today()
string += "System: %s" % platform.platform()

# standard code to display the string
to = self.GetTableOutput()
arr = vtk.vtkStringArray()
arr.SetName("Text")
arr.SetNumberOfComponents(1)
arr.InsertNextValue(string)
to.AddColumn(arr)

```

2.4.5 Python Algorithm

Programmable Source and Programmable Filter are convenient ways to prototype a Python-based data processing module. If you want to distribute such modules, or package them into modules with user interfaces, for example, then `VTKPythonAlgorithmBase`-based approach is recommended instead. Here, you write a Python class by subclassing `VTKPythonAlgorithmBase` and implementing methods to do the data processing, just like any other VTK-based filter or source. Using Python syntactic add-ons called *decorators* to annotate your class, you can easily expose the class to ParaView as a filter or a source, and ParaView will automatically add UI widgets to control parameters, etc.

Let's start with the simple script featured in Fig. 2.18. Here's the Python script to create a `VTKPythonAlgorithmBase` subclass for the same operation.

```
from vtkmodules.vtkCommonDataModel import vtkDataSet
from vtkmodules.util.vtkAlgorithm import VTKPythonAlgorithmBase
from vtkmodules.numpy_interface import dataset_adapter as dsa

class HalfVFilter(VTKPythonAlgorithmBase):
    def __init__(self):
        VTKPythonAlgorithmBase.__init__(self)

    def RequestData(self, request, inInfo, outInfo):
        # get the first input.
        input0 = dsa.WrapDataObject(vtkDataSet.GetData(inInfo[0]))

        # compute a value.
        data = input0.PointData["V"] / 2.0

        # add to output
        output = dsa.WrapDataObject(vtkDataSet.GetData(outInfo))
        output.PointData.append(data, "V_half");
        return 1
```

To expose this filter in ParaView, you have to add decorators to the class definition as follows:

```
# same imports as earlier.
from vtkmodules.vtkCommonDataModel import vtkDataSet
from vtkmodules.util.vtkAlgorithm import VTKPythonAlgorithmBase
from vtkmodules.numpy_interface import dataset_adapter as dsa

# new module for ParaView-specific decorators.
from paraview.util.vtkAlgorithm import smproxy, smproperty, smdomain

@smproxy.filter(label="Half-V Filter")
@smproperty.input(name="Input")
class HalfVFilter(VTKPythonAlgorithmBase):
    # the rest of the code here is unchanged.
    def __init__(self):
        VTKPythonAlgorithmBase.__init__(self)

    def RequestData(self, request, inInfo, outInfo):
        # get the first input.
        input0 = dsa.WrapDataObject(vtkDataSet.GetData(inInfo[0]))

        # compute a value.
```

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```

data = input0.PointData["V"] / 2.0

# add to output
output = dsa.WrapDataObject(vtkDataSet.GetData(outInfo))
output.PointData.append(data, "V_half");
return 1

```

To use this new filter, save this to a *.py file, and load it as a plugin using the **Plugin Manager** from *Tools > Plugin Manager*. On success, you will see a **Half-V Filter** in the *Filters* menu.

Besides exposing class as filters, or sources, you can use the decorators to add UI widgets to call methods on the class to set parameters.

The follow examples adds a new source named **Python-based Superquadric Source Example**, with UI to control various parameters.

```

# to add a source, instead of a filter, use the `smproxy.source` decorator.
@smproxy.source(label="Python-based Superquadric Source Example")
class PythonSuperquadricSource(VTKPythonAlgorithmBase):
    """This is dummy VTKPythonAlgorithmBase subclass that
    simply puts out a Superquadric poly data using a vtkSuperquadricSource
    internally"""
    def __init__(self):
        VTKPythonAlgorithmBase.__init__(self,
            nInputPorts=0,
            nOutputPorts=1,
            outputType='vtkPolyData')
        from vtkmodules.vtkFiltersSources import vtkSuperquadricSource
        self._realAlgorithm = vtkSuperquadricSource()

    def RequestData(self, request, inInfo, outInfo):
        from vtkmodules.vtkCommonDataModel import vtkPolyData
        self._realAlgorithm.Update()
        output = vtkPolyData.GetData(outInfo, 0)
        output.ShallowCopy(self._realAlgorithm.GetOutput())
        return 1

    # for anything too complex or not yet supported, you can explicitly
    # provide the XML for the method.
    @smproperty.xml("""
        <DoubleVectorProperty name="Center"
            number_of_elements="3"
            default_values="0 0 0"
            command="SetCenter">
            <DoubleRangeDomain name="range" />
            <Documentation>Set center of the superquadric</Documentation>
        </DoubleVectorProperty>""")
    def SetCenter(self, x, y, z):
        self._realAlgorithm.SetCenter(x,y,z)
        self.Modified()

    # In most cases, one can simply use available decorators.
    @smproperty.doublevector(name="Scale", default_values=[1, 1, 1])

```

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```

@smdomain.doublerange()
def SetScale(self, x, y, z):
    self._realAlgorithm.SetScale(x,y,z)
    self.Modified()

@smproperty.intvector(name="ThetaResolution", default_values=16)
def SetThetaResolution(self, x):
    self._realAlgorithm.SetThetaResolution(x)
    self.Modified()

@smproperty.intvector(name="PhiResolution", default_values=16)
@smdomain.intrange(min=0, max=1000)
def SetPhiResolution(self, x):
    self._realAlgorithm.SetPhiResolution(x)
    self.Modified()

@smproperty.doublevector(name="Thickness", default_values=0.3333)
@smdomain.doublerange(min=1e-24, max=1.0)
def SetThickness(self, x):
    self._realAlgorithm.SetThickness(x)
    self.Modified()

```

On loading this script as a plugin and creating the Python-based Superquadric Source Example source, the Properties panel will be populated as shown in Fig. 2.21.

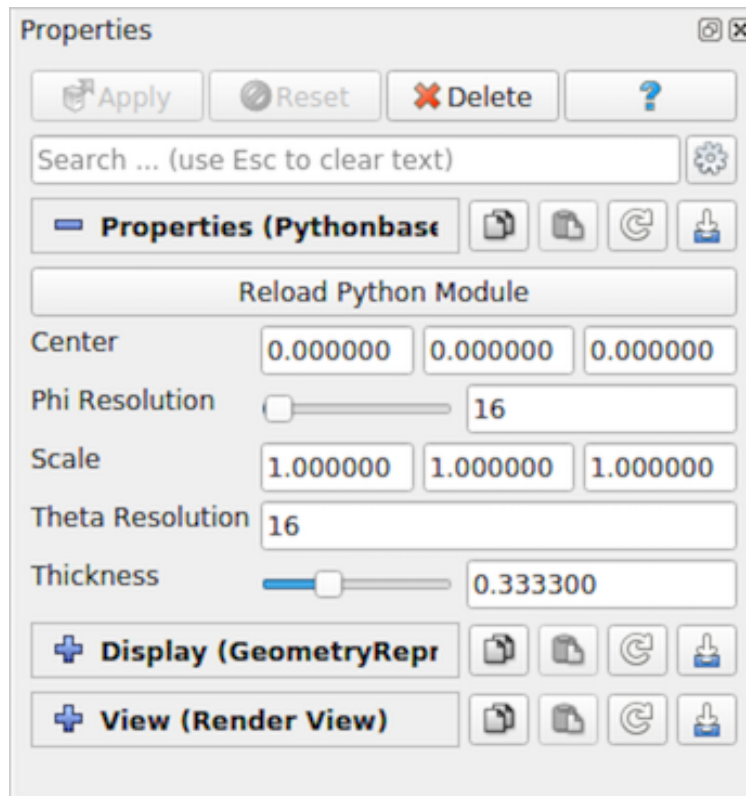


Fig. 2.21: Properties panel automatically generated from a decorated Python class, `PythonSuperquadricSource`

The decorators also enable us to add new readers and writers. Here is an example writer that uses NumPy to write tables as compressed binary arrays.

```
# `smproxy.writer` decorator register the module as writer for the provided file
# extension.
@smproxy.writer(extensions="npz", file_description="NumPy Compressed Arrays", support_
    ↴_reload=False)
@smproperty.input(name="Input", port_index=0)
# this domain lets ParaView know which types of data this writer can write.
@smdomain.datatype(dataTypes=["vtkTable"], composite_data_supported=False)
class NumpyWriter(VTKPythonAlgorithmBase):
    def __init__(self):
        VTKPythonAlgorithmBase.__init__(self, nInputPorts=1, nOutputPorts=0, inputType=
            ↴'vtkTable')
        self._filename = None

    @smproperty.stringvector(name="FileName", panel_visibility="never")
    @smdomain.filelist()
    def SetFileName(self, fname):
        """Specify filename for the file to write."""
        if self._filename != fname:
            self._filename = fname
            self.Modified()

    def RequestData(self, request, inInfoVec, outInfoVec):
        from vtkmodules.vtkCommonDataModel import vtkTable
        from vtkmodules.numpy_interface import dataset_adapter as dsa

        table = dsa.WrapDataObject(vtkTable.GetData(inInfoVec[0], 0))
        kwargs = {}
        for fname in table.RowData.keys():
            kwargs[fname] = table.RowData[fname]

        import numpy
        numpy.savez_compressed(self._filename, **kwargs)
        return 1

    def Write(self):
        self.Modified()
        self.Update()
```

2.5 Using NumPy for processing data

In Section 2.4, we looked at several recipes for writing Python script for data processing that relied heavily on using NumPy for accessing arrays and performing operations on them. In this chapter, we take a closer look at the VTK-NumPy integration layer that makes it possible to use VTK and NumPy together, despite significant differences in the data representations between the two systems.

2.5.1 Teaser

Let's start with a teaser by creating a simple pipeline with Sphere connected to an Elevation filter, followed by the Programmable Filter . Let's see how we would access the input data object in the Script for the Programmable Filter .

```
from paraview.vtk.numpy_interface import dataset_adapter as dsa
from paraview.vtk.numpy_interface import algorithms as algs

data = inputs[0]
print(data.PointData.keys())
print(data.PointData['Elevation'])
```

This example prints out the following in the output window.

```
['Normals', 'Elevation']
[ 0.67235619  0.32764378  0.72819519  0.7388373   0.70217478  0.62546903
  0.52391261  0.41762003  0.75839448  0.79325461  0.77003199  0.69332629
  0.57832992  0.44781935  0.72819519  0.7388373   0.70217478  0.62546903
  0.52391261  0.41762003  0.65528756  0.60746235  0.53835285  0.46164712
  0.39253765  0.34471244  0.58238    0.47608739  0.37453097  0.29782525
  0.2611627   0.27180481  0.55218065  0.42167011  0.30667371  0.22996798
  0.20674542  0.24160551  0.58238    0.47608739  0.37453097  0.29782525
  0.2611627   0.27180481  0.65528756  0.60746235  0.53835285  0.46164712
  0.39253765  0.34471244]
```

The importance lies in the last three lines. In particular, note how we used a different API to access the PointData and the Elevation array in the last two lines. Also note that, when we printed the Elevation array, the output didn't look like one from a vtkdataArray. In fact:

```
elevation = data.PointData['Elevation']
print(type(elevation))

import numpy
print(isinstance(elevation, numpy.ndarray))
```

This will produce the following:

```
<class 'paraview.vtk.numpy_interface.dataset_adapter.VTKArray'>
True
```

So, a VTK array is a NumPy array? What kind of trickery is this, you say? What kind of magic makes the following possible?

```
data.PointData.append(elevation + 1, 'e plus one')
print(algs.max(elevation))
print(algs.max(data.PointData['e plus one']))
print(data.VTKObject)
```

The output here is:

```
0.7932546138763428
1.7932546138763428
vtkPolyData (0x7fa20d011c60)
```

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```
...
Point Data:
...
Number Of Arrays: 3
Array 0 name = Normals
Array 1 name = Elevation
Array 2 name = e plus one
```

It is all in the `numpy_interface` module. It ties VTK datasets and data arrays to NumPy arrays and introduces a number of algorithms that can work on these objects. There is quite a bit to this module, and we will introduce it piece by piece in the rest of this chapter.

Let's wrap up this section with one final teaser.

```
print(algs.gradient(data.PointData['Elevation']))
```

Output:

```
[[ 0.32640398  0.32640398  0.01982867]
 [ 0.32640402  0.32640402  0.01982871]
 ...
 [ 0.41252578  0.20134845  0.2212007 ]
 [ 0.41105482  0.21514832  0.0782456 ]]
```

Please note that this example is not very easily replicated by using pure NumPy. The gradient function returns the gradient of an unstructured grid – a concept that does not exist in NumPy. However, the ease-of-use of NumPy is there.

2.5.2 Understanding the dataset_adapter module

In this section, let's take a closer look at the `dataset_adapter` module. This module was designed to simplify accessing VTK datasets and arrays from Python and to provide a NumPy-style interface.

Let's continue with the example from the previous section. Remember, this script is being put in the `Programmable Filter`'s `Script`, connected to the `Sphere`, followed by the `Elevation` filter pipeline.

```
from vtk.numpy_interface import dataset_adapter as dsa
...
print(data)
print(isinstance(data, dsa.VTKObjectWrapper))
```

This will print:

```
<paraview.vtk.numpy_interface.dataset_adapter.PolyData object at 0x14b7caa50>
True
```

We can access the underlying VTK object using the `VTKObject` member:

```
print(type(data.VTKObject))
```

which produces:

```
<type 'vtkCommonDataModelPython.vtkPolyData'>
```

What we get as the \$inputs\$ in the `Programmable Filter` is actually a Python object that wraps the VTK data object itself. The `Programmable Filter` does this by manually calling the `WrapDataObject` function from the `vtk.numpy_interface.dataset_adapter` module on the VTK data object. Note that the `WrapDataObject` function will return an appropriate wrapper class for all `vtkDataSet` subclasses, `vtkTable`, and all `vtkCompositeData` subclasses. Other `vtkDataObject` subclasses are not currently supported.

`VTKObjectWrapper` forwards VTK methods to its `VTKObject` so the VTK API can be accessed directly as follows:

```
print(data.GetNumberOfCells())
96L
```

However, `VTKObjectWrapper`s cannot be directly passed to VTK methods as an argument.

```
from paraview.vtk.vtkFiltersGeneral import vtkShrinkPolyData
s = vtkShrinkPolyData()
s.SetInputData(data)
```

This attempt to set the data results in an error message.

```
TypeError: SetInputData argument 1: method requires a VTK object
```

Instead, we must pass the VTK object to the VTK filter, like so:

```
s.SetInputData(data.VTKObject)
```

An important thing to note in the example above is how the Python class `vtkShrinkPolyData` was imported for use in the script. In VTK, classes are organized into different groups of related functionality, and these groups can be individually imported as Python modules. To use a class, you first identify the module in which it resides, which can be determined from the Doxygen documentation of the class [KitwareInc]. Go to the Doxygen page for the class, find the path of the file from which the documentation was generated at the bottom of the page, which has the form `dox/<first directory>/<second directory>/<class name>`. The module is then derived as `vtk<first directory><second directory>`. As an example, the documentation for `vtkShrinkPolyData` is generated from `dox/Filters/General/vtkShrinkPolyData`, hence its module is `vtkFiltersGeneral`. Then, you can import the class with a statement of the form.

Dataset attributes

So far, we have a wrapper for VTK data objects that partially behaves like a VTK data object. This gets a little bit more interesting when we start looking at how to access the fields (arrays) contained within this dataset.

For simplicity, we will embed the output generated by the script in the code itself and use the `>>>` prefix to differentiate the code from the output.

```
>>> print(data.PointData)
<vtk.numpy_interface.dataset_adapter.DataSetAttributes at 0x110f5b750>

>>> print(data.PointData.keys())
['Normals', 'Elevation']

>>> print(data.CellData.keys())
[]

>>> print(data.PointData['Elevation'])
VTKArray([ 0.5        ,  0.          ,  0.45048442,  0.3117449 ,  0.11126047,
```

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```

0.        ,  0.        ,  0.        ,  0.45048442,  0.3117449 ,
0.11126047,  0.        ,  0.        ,  0.        ,  0.45048442,
...,
0.11126047,  0.        ,  0.        ,  0.        ,  0.45048442,
0.3117449 ,  0.11126047,  0.        ,  0.        ,  0.        ], dtype=float32)

>>> elevation = data.PointData['Elevation']

>>> print(elevation[:5])
VTKArray([0.5, 0., 0.45048442, 0.3117449, 0.11126047], dtype=float32)
# Note that this works with composite datasets as well:

>>> mb = vtk.vtkMultiBlockDataSet()
>>> mb.SetNumberOfBlocks(2)
>>> mb.SetBlock(0, data.VTKObject)
>>> mb.SetBlock(1, data.VTKObject)
>>> mbw = dsa.WrapDataObject(mb)
>>> print(mbw.PointData)
<vtk.numpy_interface.dataset_adapter.CompositeDataSetAttributes instance at 0x11109f758>

>>> print(mbw.PointData.keys())
['Normals', 'Elevation']

>>> print(mbw.PointData['Elevation'])
<vtk.numpy_interface.dataset_adapter.VTKCompositeDataArray at 0x1110a32d0>

```

It is possible to access `PointData` , `CellData` , `FieldData` , `Points` (subclasses of `vtkPointSet` only), and `Polygons` (`vtkPolyData` only) this way. We will continue to add accessors to more types of arrays through this API.

2.5.3 Working with arrays

For this section, let's change our test pipeline to consist of the Wavelet source connected to the Programmable Filter .

In the Script , we access the RTData point data array as follows:

```

# Code for 'Script'
from paraview.vtk.vtkFiltersGeneral import vtkDataSetTriangleFilter
image = inputs[0]
rtdata = image.PointData['RTData']

# Let's transform this data as well, using another VTK filter.
tets = vtkDataSetTriangleFilter()
tets.SetInputDataObject(image.VTKObject)
tets.Update()

# Here, now we need to explicitly wrap the output dataset to get a
# VTKObjectWrapper instance.
ugrid = dsa.WrapDataObject(tets.GetOutput())
rtdata2 = ugrid.PointData['RTData']

```

Here, we created two datasets: an image data (`vtkImageData`) and an unstructured grid (`vtkUnstructuredGrid`). They essentially represent the same data but the unstructured grid is created by tetrahedralizing the image data. So, we expect the unstructured grid to have the same points but more cells (tetrahedra).

```
from paraview.vtk.vtkFiltersGeneral import vtkDataSetTriangleFilter
```

The array API

`numpy_interface` array objects behave very similar to NumPy arrays. In fact, arrays from `vtkDataSet` subclasses are instances of `VTKArray`, which is a subclass of `numpy.ndarray`. Arrays from `vtkCompositeDataSet` and subclasses are not NumPy arrays, but they behave very similarly. We will outline the differences in a separate section. Let's start with the basics. All of the following work as expected.

As before, for simplicity, we will embed the output generated by the script in the code itself and use the `>>>` prefix to differentiate the code from the output.

```
>>> print(rtdata[0])
60.763466

>>> print(rtdata[-1])
57.113735

>>> print(repr(rtdata[0:10:3]))
VTKArray([ 60.76346588,  95.53707886,  94.97672272, 108.49817657], dtype=float32)

>>> print(repr(rtdata + 1))
VTKArray([ 61.76346588,  86.87795258,  73.80931091, ...,  68.51051331,
          44.34006882,  58.1137352 ], dtype=float32)

>>> print(repr(rtdata < 70))
VTKArray([ True , False, False, ...,  True,  True,  True])

# We will cover algorithms later. This is to generate a vector field.
>>> avector = algs.gradient(rtdata)

# To demonstrate that avector is really a vector
>>> print(algs.shape(rtdata))
(9261,)

>>> print(algs.shape(avector))
(9261, 3)

>>> print(repr(avector[:, 0]))
VTKArray([ 25.69367027,  6.59600449,  5.38400745, ..., -6.58120966,
          -5.77147198, 13.19447994])
```

A few things to note in this example:

- Single component arrays always have the following shape: (n-tuples,) and not (n-tuples, 1)
- Multiple component arrays have the following shape: (n-tuples, n-components)
- Tensor arrays have the following shape: (n-tuples, 3, 3)
- The above holds even for images and other structured data. All arrays have one dimension (1 component arrays), two dimensions (multi-component arrays), or three dimensions (tensor arrays).

One more cool thing: It is possible to use boolean arrays to index arrays. Thus, the following works very nicely:

```
>>> print(repr(rtdata[rtdata < 70]))
VTKArray([ 60.76346588,  66.75043488,  69.19681549,  50.62128448,
          64.8801651 ,  57.72655106,  49.75050354,  65.05570221,
          57.38450241,  69.51113129,  64.24596405,  67.54656982,
          ...,
          61.18143463,  66.61872864,  55.39360428,  67.51051331,
          43.34006882,  57.1137352 ], dtype=float32)

>>> print(repr(avector[avector[:,0] > 10]))
VTKArray([[ 25.69367027,   9.01253319,   7.51076698],
          [ 13.1944809 ,   9.01253128,   7.51076508],
          [ 25.98717642,  -4.49800825,   7.80427408],
          ...,
          [ 12.9009738 ,  -16.86548471,  -7.80427504],
          [ 25.69366837,  -3.48665428,  -7.51076889],
          [ 13.19447994,  -3.48665524,  -7.51076794]])
```

Algorithms

You can do a lot simply using the array API. However, things get much more interesting when we start using the `numpy_interface.algorithms` module. We introduced it briefly in the previous examples. We will expand on it a bit more here. For a full list of algorithms, use `help(algs)`. Here are some self-explanatory examples:

```
>>> import paraview.vtk.numpy_interface.algorithms as algs
>>> print(repr(algs.sin(rtdata)))
VTKArray([-0.87873501, -0.86987603, -0.52497    , ..., -0.99943125,
          -0.59898132,   0.53547275], dtype=float32)

>>> print(repr(algs.min(rtdata)))
VTKArray(37.35310363769531)

>>> print(repr(algs.max(avector)))
VTKArray(34.781060218811035)

>>> print(repr(algs.max(avector, axis=0)))
VTKArray([ 34.78106022,  29.01940918,  18.34743023])

>>> print(repr(algs.max(avector, axis=1)))
VTKArray([ 25.69367027,   9.30603981,   9.88350773, ...,  -4.35762835,
          -3.78016186,  13.19447994])
```

If you haven't used the `axis` argument before, it is pretty easy. When you don't pass an `axis` value, the function is applied to all values of an array without any consideration for dimensionality. When `axis=0`, the function will be applied to each component of the array independently. When `axis=1`, the function will be applied to each tuple independently. Experiment if this is not clear to you. Functions that work this way include `sum`, `min`, `max`, `std`, and `var`.

Another interesting and useful function is where the indices of an array are returned where a particular condition occurs.

```
>>> print(repr(algs.where(rtdata < 40)))
(array([ 420, 9240]),)
# For vectors, this will also return the component index if an axis is not
```

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```
# defined.

>>> print(repr(algs.where(avector < -29.7)))
(VTKArray([4357, 4797, 4798, 4799, 5239]), VTKArray([1, 1, 1, 1, 1]))
```

So far, all of the functions that we discussed are directly provided by NumPy. Many of the NumPy ufuncs are included in the algorithms module. They all work with single arrays and composite data arrays. Algorithms also provide some functions that behave somewhat differently than their NumPy counterparts. These include cross, dot, inverse, determinant, eigenvalue, eigenvector, etc. See a non-exhaustive list in [Section 1.5.9](#). All of these functions are applied to each tuple rather than to a whole array/matrix. For example:

```
>>> amatrix = algs.gradient(avector)
>>> print(repr(algs.determinant(amatrix)))
VTKArray([-1221.2732624, -648.48272183, -3.55133937, ..., 28.2577152,
-629.28507693, -1205.81370163])
```

Note that everything above only leveraged per-tuple information and did not rely on the mesh. One of VTK's biggest strengths is that its data model supports a large variety of meshes, while its algorithms work generically on all of these mesh types. The algorithms module exposes some of this functionality. Other functions can be easily implemented by leveraging existing VTK filters. We used gradient before to generate a vector and a matrix. Here it is again:

```
>>> avector = algs.gradient(rtdata)
>>> amatrix = algs.gradient(avector)
```

Functions like this require access to the dataset containing the array and the associated mesh. This is one of the reasons why we use a subclass of ndarray in dataset_adapter:

```
>>> print(repr(rtdata.DataSet))
<paraview.vtk.numpy_interface.dataset_adapter.DataSet at 0x11b61e9d0>
```

Each array points to the dataset containing it. Functions such as gradient use the mesh and the array together. NumPy provides a gradient function too, you say. What is so exciting about yours? Well, this:

```
>>> print(repr(algs.gradient(rtdata2)))
VTKArray([[ 25.46767712,  8.78654003,  7.28477383],
[ 6.02292252,  8.99845123,  7.49668884],
[ 5.23528767,  9.80230141,  8.3005352 ],
...,
[ -6.43249083, -4.27642155, -8.30053616],
[ -5.19838905, -3.47257614, -7.49668884],
[ 13.42047501, -3.26066017, -7.28477287]])
>>> print(rtdata2.DataSet.GetClassName())
vtkUnstructuredGrid
```

Gradient and algorithms that require access to a mesh work whether that mesh is a uniform grid, a curvilinear grid, or an unstructured grid thanks to VTK's data model. Take a look at various functions in the algorithms module to see all the cool things that can be accomplished using it. In the remaining sections, we demonstrate how specific problems can be solved using these modules.

2.5.4 Handling composite datasets

In this section, we take a closer look at composite datasets. For this example, our pipeline is Sphere source, and Cone source is set as two inputs to the Programmable Filter .

We can create a multiblock dataset in the Programmable Filter 's Script as follows:

```
# Let's assume inputs[0] is the output from Sphere and
# inputs[1] is the output from Cone.
mb = vtk.vtkMultiBlockDataSet()
mb.SetBlock(0, inputs[0].VTKObject)
mb.SetBlock(1, inputs[1].VTKObject)
```

Many of VTK's algorithms work with composite datasets without any change. For example:

```
e = vtk.vtkElevationFilter()
e.SetInputData(mb)
e.Update()

mbe = e.GetOutputDataObject(0)
print(mbe.GetClassName())
```

This will output `vtkMultiBlockDataSet`.

Now that we have a composite dataset with a scalar, we can use `numpy_interface` . As before, for simplicity, we will embed the output generated by the script in the code itself and use the `>>>` prefix to differentiate the code from the output.

```
>>> from paraview.vtk.numpy_interface import dataset_adapter as dsa
>>> mbw = dsa.WrapDataObject(mbe)
>>> print(repr(mbw.PointData.keys()))
['Normals', 'Elevation']
>>> elev = mbw.PointData['Elevation']
>>> print(repr(elev))
<paraview.vtk.numpy_interface.dataset_adapter.VTKComposite dataArray at 0x1189ee410>
```

Note that the array type is different than we have previously seen (`VTKArray`). However, it still works the same way.

```
>>> from paraview.vtk.numpy_interface import algorithms as algs
>>> print(algs.max(elev))
0.5
>>> print(algs.max(elev + 1))
1.5
```

You can individually access the arrays of each block as follows.

```
>>> print(repr(elev.Arrays[0]))
VTKArray([ 0.5        ,  0.         ,  0.45048442,  0.3117449 ,  0.11126047,
          0.         ,  0.         ,  0.         ,  0.45048442,  0.3117449 ,
          0.11126047,  0.         ,  0.         ,  0.         ,  0.45048442,
          0.3117449 ,  0.11126047,  0.         ,  0.         ,  0.         ,
          0.45048442,  0.3117449 ,  0.11126047,  0.         ,  0.         ,
          0.         ,  0.45048442,  0.3117449 ,  0.11126047,  0.         ,
          0.         ,  0.         ,  0.45048442,  0.3117449 ,  0.11126047,
          0.         ,  0.         ,  0.45048442,  0.3117449 ,  0.11126047,
```

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```
0.11126047,  0.          ,  0.          ,  0.          ,  0.45048442,
0.3117449 ,  0.11126047,  0.          ,  0.          ,  0.          ], dtype=float32)
```

Note that indexing is slightly different.

```
>>> print(elev[0:3])
[VTKArray([ 0.5,  0.,  0.45048442], dtype=float32),
 VTKArray([ 0.,  0.,  0.43301269], dtype=float32)]
```

The return value is a composite array consisting of two VTKArrays. The [] operator simply returned the first four values of each array. In general, all indexing operations apply to each VTKArray in the composite array collection. It is similar for algorithms, where:

```
>>> print(algs.where(elev < 0.5))
[(array([ 1,  2,  3,  4,  5,  6,  7,  8,  9, 10, 11, 12, 13, 14, 15, 16, 17,
        18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34,
        35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49]),),
 (array([0, 1, 2, 3, 4, 5, 6]),)]
```

Now, let's look at the other array called Normals.

```
>>> normals = mbw.PointData['Normals']
>>> print(repr(normals.Arrays[0]))
VTKArray([[ 0.00000000e+00,  0.00000000e+00,  1.00000000e+00],
           [ 0.00000000e+00,  0.00000000e+00, -1.00000000e+00],
           [ 4.33883727e-01,  0.00000000e+00,  9.00968850e-01],
           [ 7.81831503e-01,  0.00000000e+00,  6.23489797e-01],
           [ 9.74927902e-01,  0.00000000e+00,  2.22520933e-01],
           ...
           [ 6.89378142e-01, -6.89378142e-01,  2.22520933e-01],
           [ 6.89378142e-01, -6.89378142e-01, -2.22520933e-01],
           [ 5.52838326e-01, -5.52838326e-01, -6.23489797e-01],
           [ 3.06802124e-01, -3.06802124e-01, -9.00968850e-01]], dtype=float32)
>>> print(repr(normals.Arrays[1]))
<paraview.vtk.numpy_interface.dataset_adapter.VTKNoneArray at 0x1189e7790>
```

Notice how the second array is a VTKNoneArray . This is because vtkConeSource does not produce normals. Where an array does not exist, we use a VTKNoneArray as placeholder. This allows us to maintain a one-to-one mapping between datasets of a composite dataset and the arrays in the VTKCompositeDataSet. It also allows us to keep algorithms working in parallel without a lot of specialized code.

Where many of the algorithms apply independently to each array in a collection, some algorithms are global. Take `min` and `max` , for example, as we demonstrated above. It is sometimes useful to get per-block answers. For this, you can use `_per_block` algorithms.

```
>>> print(algs.max_per_block(elev))
[0.5, 0.4330127]
```

These work very nicely together with other operations. For example, here is how we can normalize the elevation values in each block.

```
>>> _min = algs.min_per_block(elev)
>>> _max = algs.max_per_block(elev)
```

(continues on next page)

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```
>>> _norm = (elev - _min) / (_max - _min)
>>> print(algs.min(_norm))
0.0
>>> print(algs.max(_norm))
1.0
```

Once you grasp these features, you should be able to use composite arrays very similarly to single arrays.

A final note on composite datasets: The composite data wrapper provided by `numpy_interface.dataset_adapter` offers a few convenience functions to traverse composite datasets. Here is a simple example:

```
>>> for ds in mbw:
...     print(type(ds))
<class 'paraview.vtk.numpy_interface.dataset_adapter.PolyData'>
<class 'paraview.vtk.numpy_interface.dataset_adapter.PolyData'>
```

2.6 Remote and parallel visualization

One of the goals of the **ParaView** application is enabling data analysis and visualization for large datasets. **ParaView** was born out of the need for visualizing simulation results from simulations run on supercomputing resources that are often too big for a single desktop machine to handle. To enable interactive visualization of such datasets, **ParaView** uses remote and/or parallel data processing. The basic concept is that if a dataset cannot fit on a desktop machine due to memory or other limitations, we can split the dataset among a cluster of machines, driven from your desktop. In this chapter, we will look at the basics of remote and parallel data processing using **ParaView**. For information on setting up clusters, please refer to the **ParaView** Wiki [[ThePCommunity](#)].

Did you know?

Remote and parallel processing are often used together, but they refer to different concepts, and it is possible to have one without the other.

In the case of ParaView, remote processing refers to the concept of having a client, typically `paraview` or `pvython`, connecting to a `pvserver`, which could be running on a different, remote machine. All the data processing and, potentially, the rendering can happen on the `pvserver`. The client drives the visualization process by building the visualization pipeline and viewing the generated results.

Parallel processing refers to a concept where instead of single core — which we call a **rank** — processing the entire dataset, we split the dataset among multiple ranks. Typically, an instance of `pvserver` runs in parallel on more than one rank. If a client is connected to a server that runs in parallel, we are using both remote and parallel processing.

In the case of `pbatch`, we have an application that operates in parallel but without a client connection. This is a case of parallel processing without remote processing.

2.6.1 Understanding remote processing

Let's consider a simple use-case. Let's say you have two computers, one located at your office and another in your home. The one at the office is a nicer, beefier machine with larger memory and computing capabilities than the one at home. That being the case, you often run your simulations on the office machine, storing the resulting files on the disk attached to your office machine. When you're at work, to visualize those results, you simply launch `paraview` and open the data file(s). Now, what if you need to do the visualization and data analysis from home? You have several options:

- You can copy the data files over to your home machine and then use `paraview` to visualize them. This is tedious, however, as you not only have to constantly keep copying/updating your files manually, but your machine has poorer performance due to the decreased compute capabilities and memory available on it!
- You can use a desktop sharing system like *Remote Desktop* or *VNC*, but those can be flaky depending on your network connection.

Alternatively, you can use **ParaView**'s remote processing capabilities. The concept is fairly simple. You have two separate processes: `pvserver` (which runs on your work machine) and a `paraview` client (which runs on your home machine). They communicate with each other over sockets (over an SHH tunnel, if needed). As far as using `paraview` in this mode, it's no different than how we have been using it so far – you create pipelines and then look at the data produced by those pipelines in views and so on. The pipelines themselves, however, are created remotely on the `pvserver` process. Thus, the pipelines have access to the disks on your work machine. The Open File dialog will in fact browse the file system on your work machine, i.e., the machine on which `pvserver` is running. Any filters that you create in your visualization pipeline execute on the `pvserver`.

While all the data processing happens on the `pvserver`, when it comes to rendering, `paraview` can be configured to either do the rendering on the server process and deliver only images to the client (remote rendering) or to deliver the geometries to be rendered to the client and let it do the rendering locally (local rendering). When remote rendering, you'll be using the graphics capabilities on your work machine (the machine running the `pvserver`). Every time a new rendering needs to be obtained (for example, when pipeline parameters are changed or you interact with the camera, etc.), the `pvserver` process will re-render a new image and deliver that to the client. When local rendering, the geometries to be rendered are delivered to the client and the client renders those locally. Thus, not all interactions require server-side processing. Only when the visualization pipeline is updated does the server need to deliver updated geometries to the client.

2.6.2 Remote visualization in paraview

Starting a remote server

To begin using **ParaView** for remote data processing and visualization, we must first start the server application `pvserver` on the remote system. To do this, connect to your remote system using a shell and run:

```
> pvserver
```

You will see this startup message on the terminal:

```
Waiting for client...
Connection URL: cs://myhost:11111
Accepting connection(s): myhost:11111
```

This means that the server has started and is listening for a connection from a client.

Configuring a server connection

To connect to this server with the paraview client, select *File > Connect* or click the  icon in the toolbar to bring up the Choose Server Configuration dialog.

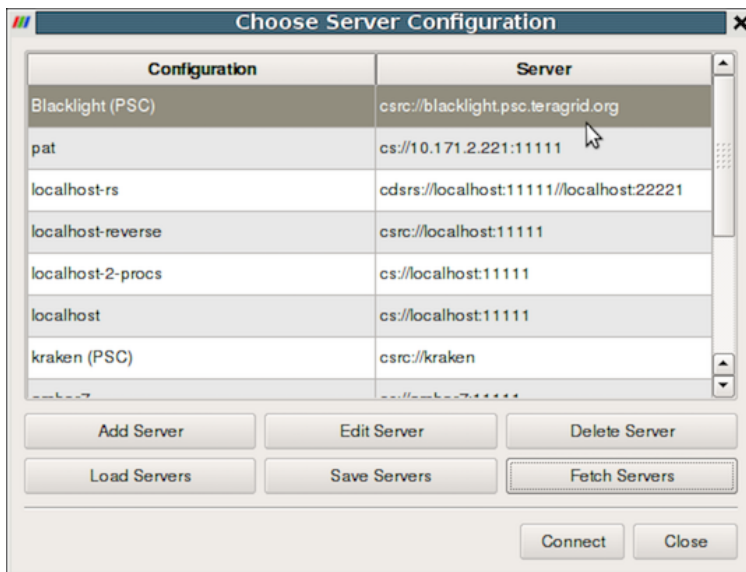


Fig. 2.22: The Choose Server Configuration dialog is used to connect to a server.

Common Errors

If your server is behind a firewall and you are attempting to connect to it from outside the firewall, the connection may not be established successfully. You may also try reverse connections ([Section 2.6.4](#)) as a workaround for firewalls. Please consult your network manager if you have network connection problems.

Figure [Fig. 2.22](#) shows the Choose Server Configuration dialog with a number of entries for remote servers. In the figure, a number of servers have already been configured, but when you first open this dialog, this list will be empty. Before you can connect to a remote server, you will need to add an entry to the list by clicking on the Add Server button. When you do, you will see the Edit Server Configuration dialog as in Figure [Fig. 2.23](#).

You will need to set a name for the connection, the server type, the DNS name of the host on which you just started the server, and the port. The default Server Type is set to Client / Server, which means that the server will be listening for an incoming connection from the client. There are several other options for this setting that we will discuss later.

When you are done, click the Configure button. Another dialog, as shown in [Fig. 2.24](#), will appear where you specify how to start the server. Since we started the server manually, we will leave the Startup Type on the default Manual setting. You can optionally set the Startup Type to Command and specify an external shell command to launch a server process.

When you click the Save button, this particular server configuration will be saved for future use. You can go back and edit the server configuration by selecting the entry in the list of servers and clicking the Edit Server button in the Choose Server Configuration dialog. You can delete it by clicking the Delete button.

Server configurations can be imported and exported through the Choose Server Configuration dialog. Use the Load Servers button to load a server configuration file and the Save Servers button to save a server configuration file. Files can be exchanged with others to access the same remote servers.

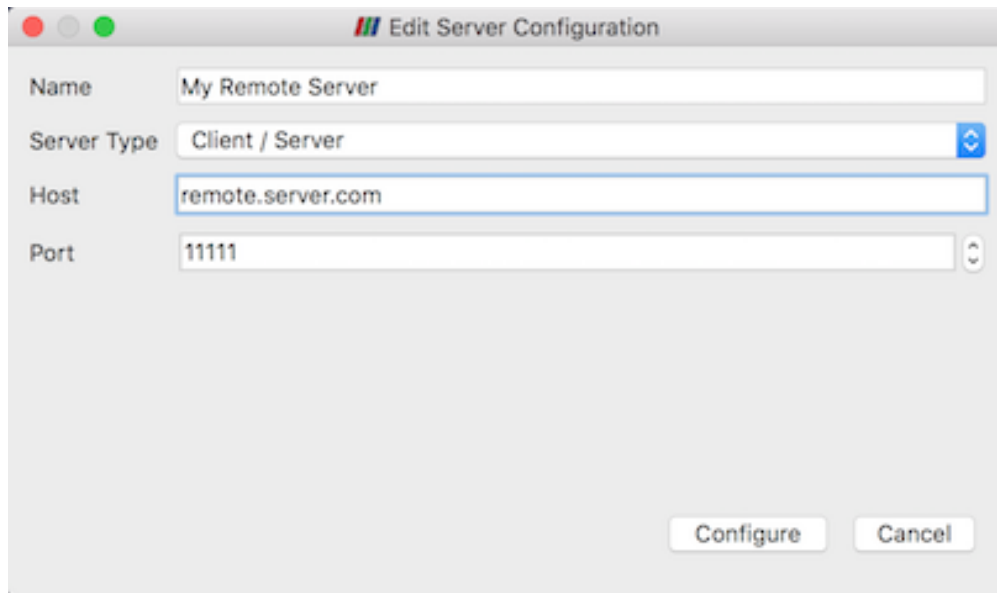


Fig. 2.23: The `Edit Server Configuration` dialog is used to configure settings for connecting to remote servers.

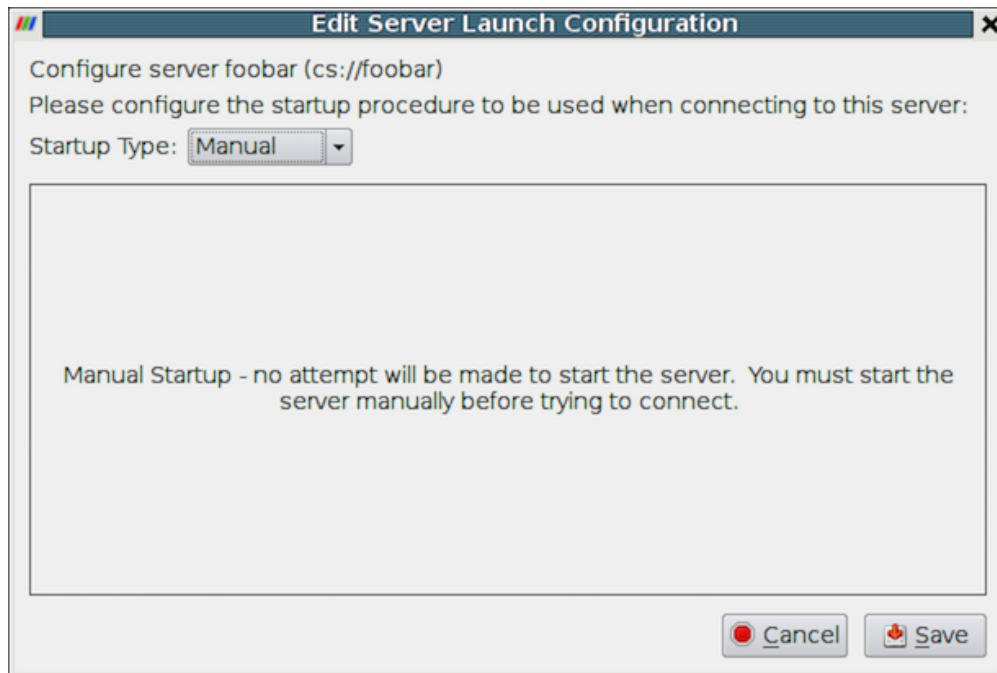


Fig. 2.24: Configure the server manually. It must be started outside of ParaView.

Did you know?

Visualization centers can provide system-wide server configurations on web servers to allow non-experts to simply select an already configured **ParaView** server. These site-wide settings can be loaded with the Fetch Servers button. Advanced users may also want to specify their own servers in more details. These features are provided thanks to ParaView Server Configuration files ([Section 2.6.5](#)).

Connect to the remote server

To connect to the server, select the server configuration you just set up from the configuration list, modify the timeout in the timeout combo box if needed and click Connect. ParaView will try to connect to the server until it succeed or timeout is reached. In that case, you can just retry as needed. Once the connection steps succeed, we are now connected and ready to build the visualization pipelines.

Common Errors

ParaView does not perform any kind of authentication when clients attempt to connect to a server. For that reason, we recommend that you do not run `pvserver` on a computing resource that is open to the outside world.

ParaView also does not encrypt data sent between the client and server. If your data is sensitive, please ensure that proper network security measures have been taken. The typical approach is to use an SSH tunnel within your server configuration files using native SSH support ([Section 2.6.5](#)).

Managing multiple clients

`pvserver` can be configured to accept connections from multiple clients at the same time. In this case only one, called the master, can interact with the pipeline. Others clients are only allowed to visualize the data. The Collaboration Panel shares information between connected clients.

To enable this mode, `pvserver` must be started with the `--multi-clients` flag:

```
pvserver --multi-clients
```

If your remote server is accessible from many users, you may want to restrict the access. This can be done with a connect id. If your client does not have the same connect-id as the server you want to connect to, you will be prompted for a connect-id. Then, if you are the master, you can change the connect-id in the Collaboration Panel.

Note that initial value for connect-id can be set by starting the `pvserver` (and respectively `paraview`) with the `--connect-id` flag, for instance:

```
pvserver --connect-id=147
```

The master client can also disable further connections in the Collaboration Panel so you can work alone, for instance. Once you are ready, you may allow other people to connect to the `pvserver` to share a visualization. This is the default feature when `pvserver` is started with `--multi-clients --disable-further-connections`.

Setting up a client/server visualization pipeline

Using paraview when connected to a remote server is not any different than when it's being used in the default stand-alone mode. The only difference, as far as the user interface goes, is that the Pipeline Browser reflects the name of the server to which you are connected. The address of the server connection next to the  icon changes from `builtin` to `cs://myhost:11111`.

Since the data processing pipelines are executing on the server side, all file I/O also happens on the server side. Hence, the Open File dialog, when opening a new data file, will browse the file system local to the `pvserver` executable and not the paraview client.

2.6.3 Remote visualization in pvpthon

The `pvpthon` executable can be used by itself for visualization of local data, but it can also act as a client that connects to a remote `pvserver`. Before creating a pipeline in `pvpthon`, use the `Connect` function:

```
# Connect to remote server "myhost" on the default port, 11111
>>> Connect("myhost") # Connect to remote server "myhost" on a
                      # specified port
>>> Connect("myhost", 11111)
```

Now, when new sources are created, the data produced by the sources will reside on the server. In the case of `pvpthon`, all data remains on the server and images are generated on the server too. Images are sent to the client for display or for saving to the local filesystem.

2.6.4 Reverse connections

It is frequently the case that remote computing resources are located behind a network firewall, making it difficult to connect a client outside the firewall to a server behind it. **ParaView** provides a way to set up a *reverse connection* that reverses the usual client server roles when establishing a connection.

To use a remote connection, two steps must be performed. First, in `paraview`, a new connection must be configured with the connection type set to reverse. To do this, open the Choose Server Configuration dialog through the `File > Connect` menu item. Add a new connection, setting the Name to `myhost (reverse)`, and select `Client / Server (reverse connection)` for Server Type. Click `Configure`. In the Edit Server Launch Configuration dialog that comes up, set the Startup Type to `Manual`. Save the configuration. Next, select this configuration and click `Connect`. A message window will appear showing that the client is awaiting a connection from the server.

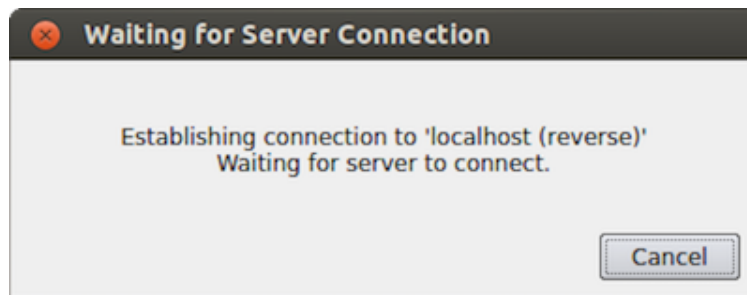


Fig. 2.25: Message window showing that the client is awaiting a connection from a server.

Second, `pvserver` must be started with the `--reverse-connection (-rc)` flag. To tell `pvserver` the name of the client, set the `--client-host (-ch)` command-line argument to the hostname of the machine on which the `paraview`

client is running. You can specify a port with the `--server-port (-sp)` command-line argument.

```
pvserver -rc --client-host=localhost --server-port=11111
```

When the server starts, it prints a message indicating the success or failure of connecting to the client. When the connection is successful, you will see the following text in the shell:

```
Connecting to client (reverse connection requested)...
Connection URL: csrv://localhost:11111
Client connected.
```

To wait for reverse connections from a `pvserver` in `pypython`, you use `ReverseConnect` instead of `Connect`.

```
# To wait for connections from a 'pvserver' on the default port 11111
>>> ReverseConnect()

# Optionally, you can specify the port number as the argument.
>>> ReverseConnect(11111)
```

2.6.5 ParaView Server Configuration Files

In the `Choose Server Configuration` dialog, it is possible to `Load Servers` and `Save Servers` using the dedicated buttons. Server configurations are stored in ParView Server Configuration files (`.pvsc`).

These files make it possible to extensively customize the server connection process. During startup, ParaView looks at several locations for server configurations to load by default.

- **On Unix-based systems and Mac OS X**

- `default_servers.pvsc` in the ParaView executable directory (you can do a `ls -l /proc/<paraview PID here>/exe` to identify the executable directory)
- `/usr/share/ParaView/servers.pvsc`
- `$HOME/.config/ParaView/servers.pvsc` (ParaView will save user defined servers here)

- **On Windows**

- `default_servers.pvsc` in the ParaView executable directory
- `%COMMON_APPDATA%\ParaView\servers.pvsc`
- `%APPDATA%\ParaView\servers.pvsc` (ParaView will save user defined servers here)

Here are a few examples of some common use-cases.

Case One: Simple command server startup

In this use-case, we are connecting to a locally started `pvserver` (`localhost`) on the `11111` port, except that the command to start the server will be automatically called just before connecting to the server, we will wait for `timeout` seconds before aborting the connection.

```
<Server name="case01" resource="cs://localhost:11111" timeout="10">
  <CommandStartup>
    <Command process_wait="0" delay="5" exec="/path/to/pvserver"/>
  </CommandStartup>
</Server>
```

Here, `CommandStartup` element specify that a command will be run before connecting to the server. The `Command` element contains the details about this command, which includes `process_wait`, the time in seconds that paraview will wait for the process to start, `delay`, the time in seconds paraview will wait after running the command to try to connect and finally, `exec`, which is the command that will be run and usually contains the path to `pvserver` but could also contain a `mpi` command to start `pvserver` distributed or to any script or executable on the `localhost` filesystem.

Case Two: Simple remote server connection

In this use-case, we are setting a configuration for a simple server connection (to a `pvserver` processes) running on a node named “`amber1`”, at port `20234`. The `pvserver` process will be started manually by the user.

```
<Server name="case02" resource="cs://amber1:20234">
  <ManualStartup/>
</Server>
```

Here, `name` specify the name of the server as it will appear in the pipeline browser, `resource` identifies the type if the connection (cs – implying client-server), host name and port. If the port number i.e. `:20234` part is not specified in the resource, then the default port number (which is `11111`) is assumed. Since the user starts `pvserver` processes manually, we use `ManualStartup`.

Case Three: Server connection with user-specified port

This is the same as case two except that we want to ask the user each time the port number to connect to the `pvserver` at.

```
<Server name="case03" resource="cs://amber1">
  <ManualStartup>
    <Options>
      <Option name="PV_SERVER_PORT" label="Server Port: ">
        <Range type="int" min="1" max="65535" step="1" default="11111" />
      </Option>
    </Options>
  </ManualStartup>
</Server>
```

Here the only difference is the `Options` element. This element is used to specify run-time options that the user specifies when connecting to the server, see this section for a list of available run-time options. In this case, we want to show the user an integral spin-box to select the port number, hence we use the `Range` element to specify the type of the option. When the user connects to this server, he is shown a dialog similarly to the following image:



Case Four: Simple connection to a data-server/render-server

This is the same as case two, except that instead of a single server (i.e. `pvserver`), we are connecting to a separate render-server/data-server with `pvdataserver` running on port 20230 on node `amber1` and `pvrenderserver` running port 20233 on node `amber2`.

```
<Server name="case04" resource="cdsrs://amber1:20230//amber2:20233">
  <ManualStartup />
</Server>
```

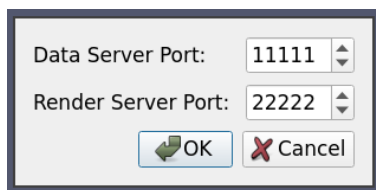
The only difference with case two, is the `resource` specification. `cdsrs` indicates that it is a client-dataserver-renderserver configuration. The first `host:port` pair is the dataserver while the second one is the render server.

Case Five: Connection to a data-server/render-server with user specified server port

This is a combination of case three and case four, where we want to ask the user for the port number for both the render server and the data server.

```
<Server name="case05" resource="cdsrs://localhost//localhost">
  <ManualStartup>
    <Options>
      <Option name="PV_DATA_SERVER_PORT" label="Data Server Port: ">
        <Range type="int" min="1" max="65535" step="1" default="11111" />
      </Option>
      <Option name="PV_RENDER_SERVER_PORT" label="Render Server Port: ">
        <Range type="int" min="1" max="65535" step="1" default="22222" />
      </Option>
    </Options>
  </ManualStartup>
</Server>
```

The XML is quite self-explanatory given what we has already been explained above. The options dialog produced by this XML looks as follows:



Case Six: Reverse Connection

By default the client connects to the server processes. However it is possible to tell the paraview client to wait for the server to connect to it instead. This is called a reverse connection. In such a case the server processes must be started with `--reverse-connection` or `--rc` flag.

To indicate reverse connection in the server configuration xml, the only change is suffixing the `resource` protocol part with `rc` (for reverse connection). eg.

```
resource="csrc://localhost" -- connect to pvserver on localhost using reverse connection
resource="cdsrsrc://localhost//localhost" -- connect to pvdataserver/pvrenderserver
  ↳ using reverse connection.
```

So a simple local reverse connection server configuration, similarly to case one, would look like this

```
<Server name="case06" resource="csrc://localhost:11111">
  <CommandStartup>
    <Command exec="/path/to/pvserver --reverse-connection --client-host=localhost"/>
  </CommandStartup>
</Server>
```

Here the --client-host=localhost in the exec is actually not needed has this is the default.

Case Seven: Server command with option

As we have seen in case one, the server can be started by ParaView on connection, but this can be combined with the Option element as seen in case three to create a dynamically generated server command.

```
<Server name="case07" resource="cs://localhost">
  <CommandStartup>
    <Options>
      <!-- The user chooses the port on which to start the server -->
      <Option name="PV_SERVER_PORT" label="Server Port: ">
        <Range type="int" min="1" max="65535" step="1" default="11111" />
      </Option>
    </Options>
    <Command delay="5" exec="/path/to/pvserver">
      <Arguments>
        <Argument value="--server-port=$PV_SERVER_PORT$" />
      </Arguments>
    </Command>
  </CommandStartup>
</Server>
```

As with case one, we are using CommandStartup and Command elements. Command line arguments can be passed to the command executed using the Arguments element. All runtime environment variables specified as \$name\$ are replaced with the actual values. Eg. in this case \$PV_SERVER_PORT\$ gets replaced by the port number chosen by the user in the options dialog.

Case Eight: Using connection-id and random port

In many cases, a server cluster may be running multiple pvserver (or pvdataserver/pvrenderserver) processes for different users. In that case we need some level of authentication between the server and the client. This can be achieved (at a very basic level) with the connect-id option. If specified on the command line when starting the server processes (using --connect-id) then the server will allow only that client which reports the same connection id to connect.

We also want to avoid port collision with other users, so we use a random port for the server connection.

Here is an example similarly to case seven but with a connect-id option and random server port.

```
<Server name="case08" resource="cs://localhost">
  <CommandStartup>
    <Options>
      <Option name="PV_CONNECT_ID" label="Connect ID" readonly="true">
        <Range type="int" min="1" max="65535" default="random" />
      </Option>
```

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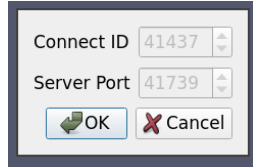
(continued from previous page)

```

<Option name="PV_SERVER_PORT" label="Server Port" readonly="true">
    <Range type="int" min="11111" max="65535" default="random" />
</Option>
</Options>
<Command exec="/path/to/pvserver" delay="5">
    <Arguments>
        <Argument value="--connect-id=$PV_CONNECT_ID$" />
        <Argument value="--server-port=$PV_SERVER_PORT$" />
    </Arguments>
</Command>
</CommandStartup>
</Server>

```

In this case, the `readonly` attribute on the `Option` indicates that the value cannot be changed by the user, it is only shown for information purposes. The default value for the `PV_CONNECT_ID` and `PV_SERVER_PORT` is set to random so that ParaView makes up a value at run time. Of course, in a production environment they should be assigned by user instead of randomly generated.



Case Nine: Starting server using ssh

In this use case the server process is spawned on some remote host using specifically crafted ssh command. We want the user to be able to specify the ssh executable. We also want to preserve the ssh executable path across ParaView sessions so that the user does not have to enter it each time.

```

<Server name="case09" resource="cs://localhost:11111">
    <CommandStartup>
        <Options>
            <Option name="SSH_USER" label="SSH Username" save="true">
                <!-- choose the username. Since 'save' is true, this value will
                    be maintained across sessions -->
                <String default="user" />
            </Option>
            <Option name="SSH_EXE" label="SSH Executable" save="true">
                <!-- select the SSH executable. Since 'save' is true, this value will
                    also be maintained across sessions -->
                <File default="ssh" />
            </Option>
        </Options>
        <Command exec="$SSH_EXE$" delay="5">
            <Arguments>
                <Argument value="-L8080:amber5:11111" /> <!-- port forwarding -->
                <Argument value="amber5" />
                <Argument value="-l" />
                <Argument value="$SSH_USER$" />
                <Argument value="/path/to/pvserver" />
            </Arguments>
        </Command>
    </CommandStartup>
</Server>

```

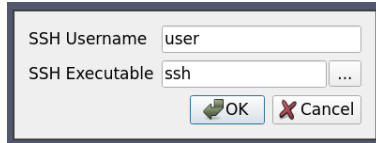
(continues on next page)

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```

</Arguments>
</Command>
</CommandStartup>
</Server>

```



Note here that the value for the `exec` attribute is set to `SSH_EXE` hence it gets replaced by the user selected ssh executable. We use the optional attribute `save` on the `Option` element to tell ParaView to preserve the user chosen value across ParaView sessions so that the user doesn't have to enter the username and the ssh executable every time he wants to connect to this server.

Did you know?

While SSH connection can be started by crafting the command, ParaView now support SSH connection natively by specifying a `SSHCommand`, see below for more information.

Case Ten: Starting server using custom script with custom user-settable options

This example will illustrate the full capability of server configuration. Suppose we have a custom script “MyServer-Starter” that takes in multiple arguments to start the server process. We want the user to be able to set up values for these arguments when he tries to connect to using this configuration. As an example, let’s say MyServerStarter takes the following arguments:

- `--force-offscreen-rendering` – to indicate use of offscreen rendering
- `--force-onscreen-rendering` – to indicate on-screen rendering (this can be assumed from absence of `--force-offscreen-rendering`, but we are using it as an example)
- `--session-name=<string>` – some string identifying the session
- `--mpitype=<mpich1.2|mpich2|openmpi>` – choose between available MPI implementations
- `--num-procs=<num>` – number of server processes
- `--server-port` – port number passed the pvserver processes

All (except the `--server-port`) of these must be settable by the user at the connection time. This can be achieved as follows:

```

<Server name="case10" resource="cs://localhost">
  <CommandStartup>
    <Options>
      <Option name="OFFSCREEN" label="Use offscreen rendering">
        <Boolean true="--use-offscreen" false="--use-onscreen" default="false" />
      </Option>
      <Option name="SESSIONID" label="Session Identifier">
        <String default="session01"/>
      </Option>
      <Option name="MPITYPE" label="MPI Implementation">

```

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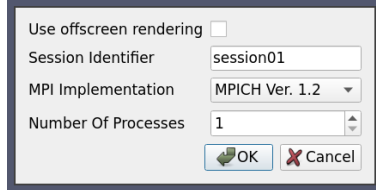
(continued from previous page)

```

<Enumeration default="mpich1.2">
    <Entry value="mpich1.2" label="MPICH Ver. 1.2" />
    <Entry value="mpich2" label="MPICH Ver 2.0" />
    <Entry value="openmpi" label="Open MPI" />
</Enumeration>
</Option>
<Option name="NUMPROC" label="Number Of Processes">
    <Range type="int" min="1" max="256" step="4" default="1" />
</Option>
</Options>
<Command exec="/path/to/MyServerStarter" delay="5">
    <Arguments>
        <Argument value="--server-port=$PV_SERVER_PORT$" />
        <Argument value="--mpitype=$MPITYPE$" />
        <Argument value="--num-procs=$NUMPROC$" />
        <Argument value="$OFFSCREEN$" />
        <Argument value="--session-name=$SESSIONID$" />
    </Arguments>
</Command>
</CommandStartup>
</Server>

```

Each Option defines a new run-time variable that can be accessed as \${name}\$ in the Command section. When the user tries to connect using this configuration, he is shown the following options dialog:



This can be extended to start the server processes using ssh or any batch scheduler etc. as may be the required by the server administrator. This can also be set up to use reverse connection (by changing the protocol in the resource attribute).

Case Eleven: Case Ten + Switch Statement

This is same as case ten with one change: We no longer allow the user to choose the number of processes. Instead, the number of processes is automatically selected based on the value of the distribution combobox.

```

<Server name="case11" resource="cs://localhost">
    <CommandStartup>
        <Options>
            <Option name="OFFSCREEN" label="Use offscreen rendering">
                <Boolean true="--use-offscreen" false="--use-onscreen" default="false" />
            </Option>
            <Option name="SESSIONID" label="Session Identifier">
                <String default="session01"/>
            </Option>
            <Option name="MPI_TYPE" label="MPI Implementation">
                <Enumeration default="mpich1.2">

```

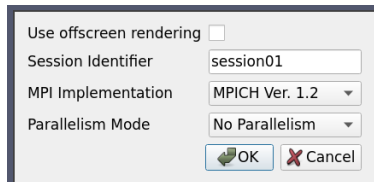
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```

<Entry value="mpich1.2" label="MPICH Ver. 1.2" />
<Entry value="mpich2" label="MPICH Ver 2.0" />
<Entry value="openmpi" label="Open MPI" />
</Enumeration>
</Option>
<Option name="DISTRIBUTION" label="Distribution Mode">
  <Enumeration default="notDistributed">
    <Entry value="notDistributed" label="Not Distributed" />
    <Entry value="someDistribution" label="Some Distribution" />
    <Entry value="highDistribution" label="Highly Distributed" />
  </Enumeration>
</Option>
<Switch name="DISTRIBUTION">
  <Case value="notDistributed">
    <Set name="NUMPROC" value="1" />
  </Case>
  <Case value="someDistribution">
    <Set name="NUMPROC" value="2" />
  </Case>
  <Case value="highDistribution">
    <Set name="NUMPROC" value="10" />
  </Case>
</Switch>
</Options>
<Command exec="/path/to/MyServerStarter" delay="5">
  <Arguments>
    <Argument value="--server-port=$PV_SERVER_PORT$" />
    <Argument value="--mpitype=$MPITYPE$" />
    <Argument value="--num-procs=$NUMPROC$" />
    <Argument value="$OFFSCREEN$" />
    <Argument value="--session-name=$SESSIONID$" />
  </Arguments>
</Command>
</CommandStartup>
</Server>

```



The `Switch` statement can only have `Case` statements as children, while the `Case` statement can only have `Set` statements as children. `Set` statements are not much different from `Option` except that the value is fixed and the user is not prompted to set that value.

Case Twelve: Simple SSH run server command

If `Command` element let you craft SSH commands, it can be quite complex to do so and the pipeline browser in ParaView may not show the correct server as it could connect through a ssh tunnel.

Here, similarly to case one, we use native ssh support to start a `pvserver` process remotely, on `amber1`, before connecting to it directly on the default port:

```
<Server name="case12" resource="cs://amber1">
  <CommandStartup>
    <SSHCommand exec="/path/to/pvserver" delay="5">
      <SSHConfig user="user"/>
    </SSHCommand>
  </CommandStartup>
</Server>
```

First `SSHCommand` element is used instead of `Command` so that ParaView knows to use native ssh support. Then the `SSHConfig` element is used to configure the ssh connection. The `user` attribute is the SSH user to use with SSH. If a password is needed, it will be asked on the terminal used to run ParaView, which may not be visible in certain cases.

Case Thirteen: SSH run server command with complex config

Here, similarly to case twelve, we use native ssh support to start a `pvserver` process remotely, on `amber1`, before connecting to it directly, but we specify much more specifically the configuration to use.

```
<Server name="case13" resource="cs://amber1">
  <CommandStartup>
    <Options>
      <!-- The user chooses the port on which to start the server -->
      <Option name="PV_SERVER_PORT" label="Server Port: ">
        <Range type="int" min="1" max="65535" step="1" default="11111" />
      </Option>
    </Options>
    <SSHCommand exec="/path/to/pvserver" delay="5">
      <SSHConfig user="user">
        <Terminal exec="/usr/bin/xterm"/>
        <SSH exec="/usr/bin/ssh"/>
      </SSHConfig>
      <Arguments>
        <Argument value="--server-port=$PV_SERVER_PORT$"/>
      </Arguments>
    </SSHCommand>
  </CommandStartup>
</Server>
```

Inside the `SSHConfig` element, we use different elements. Here, `Terminal` element is used to specify that ParaView will try to open a terminal to ask the user for his password. Here, the terminal executable is specified using the `exec` attribute. If it was not, ParaView would try to find one automatically (Linux and Windows). When troubleshooting server configuration, not using `Terminal` element is suggested as the terminal will close as soon as the command finish executing. On Linux, it is also possible to replace the `Terminal` element by the `AskPass` element to specify the ParaView should use `SSH_ASKPASS` so that a ask-pass binary is used when asking for the SSH password. Finally, the `SSH` element specify the SSH binary to use thanks to its `exec` attribute.

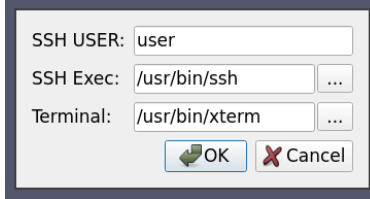
We also use `PV_SERVER_PORT`, similarly to case seven to let user select the port to connect to.

Case Fourteen: SSH run server command with user chosen config

Here, similarly to case thirteen and five, we use native ssh support to start a pvserver process remotely, on amber1, before connecting to it directly, but we let the user choose interactively some SSH options.

```
<Server name="case14" resource="cs://amber1">
  <CommandStartup>
    <Options>
      <Option label="SSH USER:" name="SSH_USER" save="true">
        <String default="user"/>
      </Option>
      <Option label="SSH Exec:" name="SSH_EXEC" save="true">
        <File default="/usr/bin/ssh" />
      </Option>
      <Option label="Terminal:" name="TERMINAL" save="true">
        <File default="/usr/bin/xterm"/>
      </Option>
    </Options>
    <SSHCommand exec="/path/to/pvserver" delay="5">
      <SSHConfig user="$SSH_USER$">
        <Terminal exec="$TERMINAL$"/>
        <SSH exec="$SSH_EXEC$"/>
      </SSHConfig>
      <Arguments>
        <Argument value="--server-port=$PV_SERVER_PORT$"/>
      </Arguments>
    </SSHCommand>
  </CommandStartup>
</Server>
```

Similarly to all other options, SSH related options can be set interactively by the user. Here we let the user set the SSH user, the SSH executable as well as the Terminal executable to use when connecting through ssh.



Case Fifteen: Ssh run server command with reverse connection

Similarly to case twelve and thirteen, we use native ssh support to start a reverse connection pvserver process remotely, on amber1, before letting it connect to ParaView using the hostname of the client on static non-default port.

```
<Server name="case15" resource="csrc://amber1:11112">
  <CommandStartup>
    <SSHCommand exec="/path/to/pvserver" delay="5">
      <SSHConfig user="user">
        <Terminal/>
      </SSHConfig>
      <Arguments>
```

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```

<Argument value="--reverse-connection"/>
<Argument value="--client-host=$PV_CLIENT_HOST$"/>
<Argument value="--server-port=$PV_SERVER_PORT$"/>
</Arguments>
</SSHCommand>
</CommandStartup>
</Server>

```

The only difference with case twelve is in the `ressource`, which now contain the reverse connection as well as the usage of `PV_CLIENT_HOST` in the arguments for the reverse connection, automatically set to the hostname of the client which the server should be able to resolve to an ip to connect to.

Case Sixteen: Secured Connection to a Server through SSH tunnel

To communicate securely through a ssh tunnel, something usually done with a crafted command looking like this: `ssh -L 8080:localhost:port user@remote /path/to/pvserver --sp=port`

You would then connect on a server on `localhost:8080` within ParaView. This is complex to set up either manually or with a `Command` element. Also, the true server and port will not appear in the pipeline browser in ParaView.

This is however natively supported with `SSHCommand` element. Here we create a secured SSH tunnel to `amber1` before connecting through the SSH tunnel on the `11111` port.

```

<Server name="case16" resource="cs://amber1:11111">
  <CommandStartup>
    <SSHCommand exec="/path/to/pvserver" delay="5">
      <SSHConfig user="user">
        <Terminal/>
        <PortForwarding local="8080"/>
      </SSHConfig>
      <Arguments>
        <Argument value="--server-port=$PV_SERVER_PORT$"/>
      </Arguments>
    </SSHCommand>
  </CommandStartup>
</Server>

```

Similarly to case thirteen, we only add a `PortForwarding` element in the `SSHConfig` element with a `local` attribute `port`, so that ParaView creates a SSH tunnel to connect through. The `PV_SERVER_PORT` is automatically set to the value of the port to use within the SSH tunnel. In ParaView, the tunnel will be integrated nicely in the UI with the correct port and hostname in the pipeline browser, the server icon will look different with a small lock to note the secured nature of this connection:



Case Seventeen: Secured Reverse Connection from a Server through SSH tunnel

Similarly to case sixteen, a reverse connection through a SSH tunnel would require to craft a command like this one:
`ssh -R 8080:localhost:port user@remote /path/to/pvserver --rc --ch=localhost --sp=8080`

You would then connect on a reverse connection server on localhost:8080 on ParaView. This is complex to set up either manually or with a `Command` element. Also, the true server and port will not appear in the pipeline browser in ParaView.

This is however natively supported with `SSHCommand`. Here we create a reverse secured SSH tunnel to `amber1` before reverse connecting through the SSH tunnel on a specific port.

```
<Server name="case17" resource="csrc://amber1:11115">
  <CommandStartup>
    <SSHCommand exec="/path/to/pvserver" delay="5">
      <SSHConfig user="user">
        <Terminal/>
        <PortForwarding local="8080"/>
      </SSHConfig>
      <Arguments>
        <Argument value="--reverse-connection"/>
        <Argument value="--client-host=localhost"/>
        <Argument value="--server-port=$PV_SSH_PF_SERVER_PORT$"/>
      </Arguments>
    </SSHCommand>
  </CommandStartup>
</Server>
```

We also specify a `PortForwarding` element in the `SSHConfig` with a local port to trigger the creation of the SSH tunnel. Finally, `$PV_SSH_PF_SERVER_PORT$` variable should be used by the server to connect through the SSH tunnel to the client.

Did you know?

While SSH native support can simplify the configuration file, some cases are still not covered and require complex custom command. Client/DataServer/RenderServer SSH setup are not supported natively, nested SSH tunnels are not supported natively either. To create such setup, use of complex `Command` is needed.

PVSC file XML Schema

Here is the exhaustive PVSC file XML schema

- The `<Servers>` tag is the root element of the document, which contains zero-to-many `<Server>` tags.
- Each `<Server>` tag represents a configured server:
 - The `name` attribute uniquely identifies the server configuration, and is displayed in the user interface.
 - The `timeout` attribute specifies the maximum amount of time (in seconds) that the client will wait for the server to start, -1 means forever, default to 60.
 - The `resource` attribute specifies the type of server connection, server host(s) and optional port(s) for making a connection. Values are

- `cs://<host>:<port>` - for client-pvserver configurations with forward connection i.e. client connects to the server. If not specified, port default to 11111.
- `csrc://<host>:<port>` - for client-pvserver configurations with reverse connection i.e. server connects to the client. If not specified, port default to 11111.
- `cdsrs://<ds-host>:<ds-port>//<rs-host>:<rs-port>` - for client-pvdataserver-pvrenderserver configurations with forward connection. If not specified, ds-port default to 11111, rs-port default to 22222.
- `cdsrsrc://<ds-host>:<ds-port>//<rs-host>:<rs-port>` - for client-pvdataserver-pvrenderserver configurations with reverse connection. If not specified, ds-port default to 11111, rs-port default to 22222.
- The `<CommandStartup>` tag is used to run an external command to start a server.
 - An optional `<Options>` tag can be used to prompt the user for options required at startup.
 - Each `<Option>` tag represents an option that the user will be prompted to modify before startup.
 - The `name` attribute defines the name of the option, which will become its variable name when used as a run-time environment variable, and for purposes of string-substitution in `<Argument>` tags.
 - The `label` attribute defines a human-readable label for the option, which will be used in the user interface.
 - The optional `readonly` attribute can be used to designate options which are user-visible, but cannot be modified.
 - The optional `save` attribute can be used to indicate that the value chosen by the user for this option will be saved in the ParaView settings so that it's preserved across ParaView sessions.
 - A `<Range>` tag designates a numeric option that is only valid over a range of values.
 - The `type` attribute controls the type of number controlled. Valid values are `int` for integers and `double` for floating-point numbers, respectively.
 - The `min` and `max` attributes specify the minimum and maximum allowable values for the option (inclusive).
 - The `step` attribute specifies the preferred amount to increment / decrement values in the user interface.
 - The `default` attribute specifies the initial value of the option.
 - As a special-case for integer ranges, a default value of `random` will generate a random number as the default each time the user is prompted for a value. This is particularly useful with `PV_CONNECT_ID` and `PV_SERVER_PORT`.
 - A `<String>` tag designates an option that accepts freeform text as its value.
 - The `default` attribute specifies the initial value of the option.
 - A `<File>` tag designates an option that accepts freeform text along with a file browse button to assist in choosing a filepath

- The **default** attribute specifies the initial value of the option.
- A **<Boolean>** tag designates an option that is either on/off or true/false.
- The **true** attribute specifies what the option value will be if enabled by the user.
- The **false** attribute specifies what the option value will be if disabled by the user.
- The **default** attribute specifies the initial value of the option, either **true** or **false**.
- An **<Enumeration>** tag designates an option that can be one of a finite set of values.
 - The **default** attribute specifies the initial value of the option, which must be one of its enumerated values.
 - Each **<Entry>** tag describes one allowed value.
 - The **name** tag specifies the value for that choice.
 - The **label** tag provides human-readable text that will be displayed in the user interface for that choice.
- A **<Command>** tag is used to specify the external command and its startup arguments.
 - The **exec** attribute specifies the filename of the command to be run. The system PATH will be used to search for the command, unless an absolute path is specified. If the value for this attribute is specified as **\$STRING\$**, then it will be replaced with the value of a predefined or user-defined (through **<Option/>**) variable.
 - The **process_wait** attribute specifies a waiting time (in seconds) that ParaView will wait for the exec command to start. Default to 0.
 - The **delay** attribute specifies a delay (in seconds) between the time the startup command completes and the time that the client attempts a connection to the server. Default to 0.
 - **<Argument>** tags are command-line arguments that will be passed to the startup command.
 - String substitution is performed on each argument, replacing each **\$STRING\$** with the value of a predefined or user-defined variable.
 - Arguments whose value is an empty string are not passed to the startup command.
- A **<SSHCommand>** tag is used to specify the external command to be started through ssh
 - All **<Command>** related attributes and tags still applies.
 - A **<SSHConfig>** tag is used to set the SSH configuration.
 - The **user** attribute is used to set the SSH username
 - A **<Terminal>** tag is used to inform ParaView to use a terminal to issue ssh commands and ask user for password when needed.

- The `exec` attribute specifies the terminal executable to use, if not set, ParaView will try to find one automatically, on Windows and Linux only.
- A `<AskPass>` tag, which should not be used with `<Terminal>` tag, can be used to inform ParaView to use a AskPass, using the `SSH_ASKPASS` environnement variable, on Linux only.
- A `<SSH>` tag, used to specify
 - the `exec` attribute that specifies the SSH executable to use.
- A `<PortForwarding>` tag, that indicates to ParaView that a SSH tunnel will need to be created, either forward or reverse depending on the connection type.
- the `local` attribute that specified the local port to use the SSH tunel.
- The `<ManualStartup>` tag indicates that the user will manually start the given server prior to connecting.
- An optional `<Options>` tag can be used to prompt the user for options required at startup. Note that `PV_SERVER_PORT`, `PV_DATA_SERVER_PORT`, `PV_RENDER_SERVER_PORT`, and `PV_CONNECT_ID` are the only variables that make sense in this context.

Startup Command Variables

When a startup command is run, its environment will include all of the user-defined variables specified in `<Option>` tags, plus the following predefined variables:

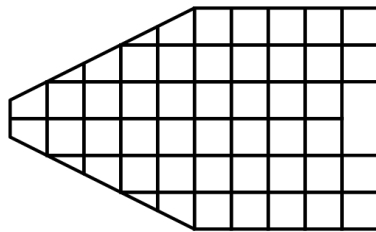
- `PV_CLIENT_HOST`
- `PV_CONNECTION_URI`
- `PV_CONNECTION_SCHEME`
- `PV_VERSION_MAJOR` (e.g. 5)
- `PV_VERSION_MINOR` (e.g. 9)
- `PV_VERSION_PATCH` (e.g. 1)
- `PV_VERSION` (e.g. 5.9)
- `PV_VERSION_FULL` (e.g. 5.9.1)
- `PV_SERVER_HOST`
- `PV_SERVER_PORT`
- `PV_SSH_PF_SERVER_PORT`
- `PV_DATA_SERVER_HOST`
- `PV_DATA_SERVER_PORT`
- `PV_RENDER_SERVER_HOST`
- `PV_RENDER_SERVER_PORT`
- `PV_CLIENT_PLATFORM` (possible values are: Windows, Apple, Linux, Unix, Unknown)
- `PV_APPLICATION_DIR`

- PV_APPLICATION_NAME
- PV_CONNECT_ID

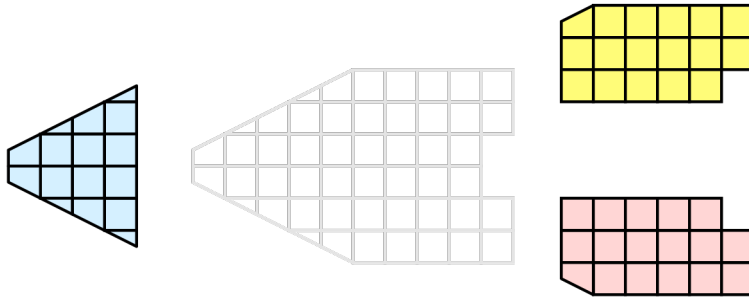
These options can be used in the <Command> or <SSHCommand> elements part of the PVSC files, as well as extracted from the environment when running the command. If an <Option> element defines a variable with the same name as a predefined variable, the <Option> element value takes precedence. This can be used to override defaults that are normally hidden from the user. As an example, if a site wants users to be able to override default port numbers, the server configuration might specify an <Option> of PV_SERVER_PORT.

2.6.6 Understanding parallel processing

Parallel processing, put simply, implies processing the data in parallel, simultaneously using multiple workers. Typically, these workers are different processes that could be running on a multicore machine or on several nodes of a cluster. Let's call these ranks. In most data processing and visualization algorithms, work is directly related to the amount of data that needs to be processed, i.e., the number of cells or points in the dataset. Thus, a straight-forward way of distributing the work among ranks is to split an input dataset into multiple chunks and then have each rank operate only an independent set of chunks. Conveniently, for most algorithms, the result obtained by splitting the dataset and processing it separately is same as the result that we'd get if we processed the dataset in a single chunk. There are, of course, exceptions. Let's try to understand this better with an example. For demonstration purposes, consider this very simplified mesh.

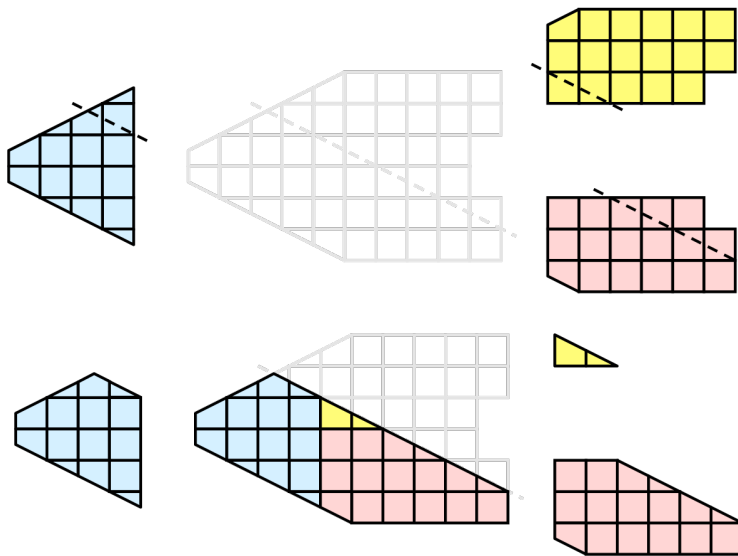


Now, let us say we want to perform visualizations on this mesh using three processes. We can divide the cells of the mesh as shown below with the blue, yellow, and pink regions.



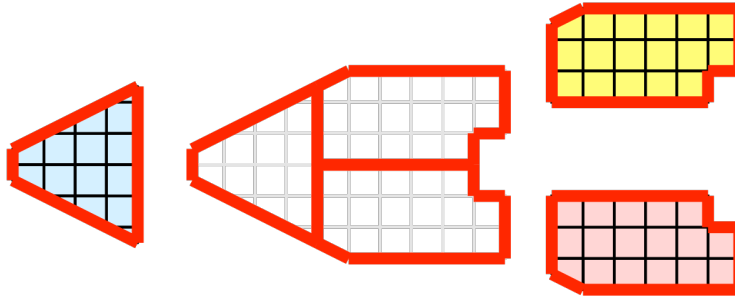
Once partitioned, some visualization algorithms will work by simply allowing each process to independently run the algorithm on its local collection of cells. Take clipping as an example. Let's say that we define a clipping plane and give that same plane to each of the processes.

Each process can independently clip its cells with this plane. The end result is the same as if we had done the clipping serially. If we were to bring the cells together (which we would never actually do for large data for obvious reasons), we would see that the clipping operation took place correctly.



Ghost levels

Unfortunately, blindly running visualization algorithms on partitions of cells does not always result in the correct answer. As a simple example, consider the **external faces** algorithm. The external faces algorithm finds all cell faces that belong to only one cell, thereby, identifying the boundaries of the mesh.

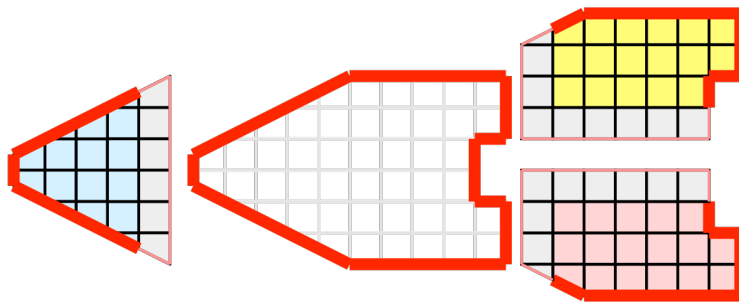


Oops! We see that when all the processes ran the external faces algorithm independently, many internal faces were incorrectly identified as being external. This happens where a cell in one partition has a neighbor in another partition. A process has no access to cells in other partitions, so there is no way of knowing that these neighboring cells exist.

The solution employed by ParaView and other parallel visualization systems is to use **ghost cells**. Ghost cells are cells that are held in one process but actually belong to another. To use ghost cells, we first have to identify all the neighboring cells in each partition. We then copy these neighboring cells to the partition and mark them as ghost cells, as indicated with the gray colored cells in the following example.

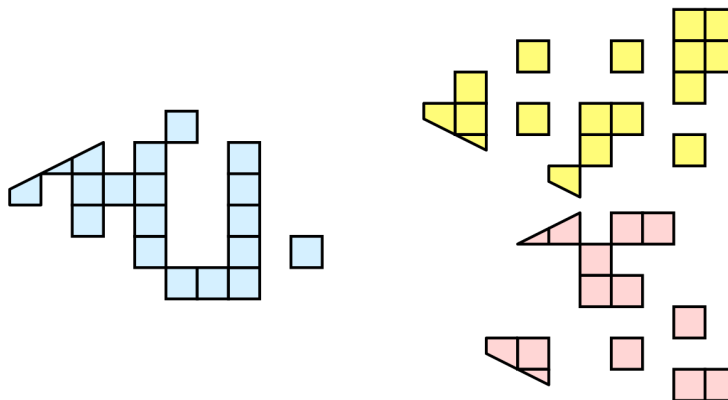
When we run the external faces algorithm with the ghost cells, we see that we are still incorrectly identifying some internal faces as external. However, all of these misclassified faces are on ghost cells, and the faces inherit the ghost status of the cell from which it came. ParaView then strips off the ghost faces, and we are left with the correct answer.

In this example, we have shown one layer of ghost cells: only those cells that are direct neighbors of the partition's cells. ParaView also has the ability to retrieve multiple layers of ghost cells, where each layer contains the neighbors of the previous layer not already contained in a lower ghost layer or in the original data itself. This is useful when we have cascading filters that each require their own layer of ghost cells. They each request an additional layer of ghost cells from upstream, and then remove a layer from the data before sending it downstream.

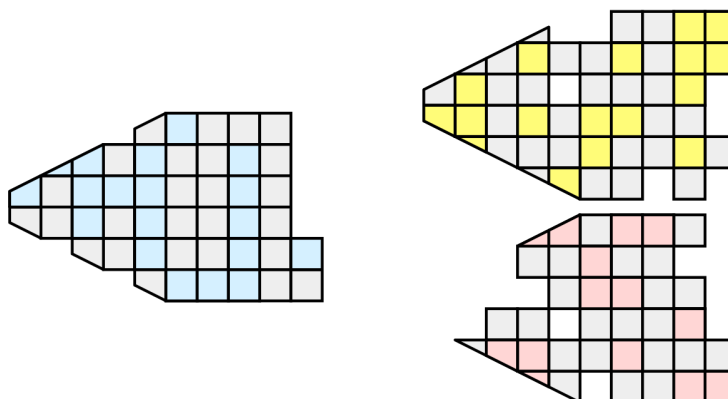


Data partitioning

Since we are breaking up and distributing our data, it is prudent to address the ramifications of how we partition the data. The data shown in the previous example has a **spatially coherent** partitioning. That is, all the cells of each partition are located in a compact region of space. There are other ways to partition data. For example, you could have a random partitioning.



Random partitioning has some nice features. It is easy to create and is friendly to load balancing. However, a serious problem exists with respect to ghost cells.



In this example, we see that a single level of ghost cells nearly replicates the entire dataset on all processes. We have thus removed any advantage we had with parallel processing. Because ghost cells are used so frequently, random partitioning is not used in ParaView.

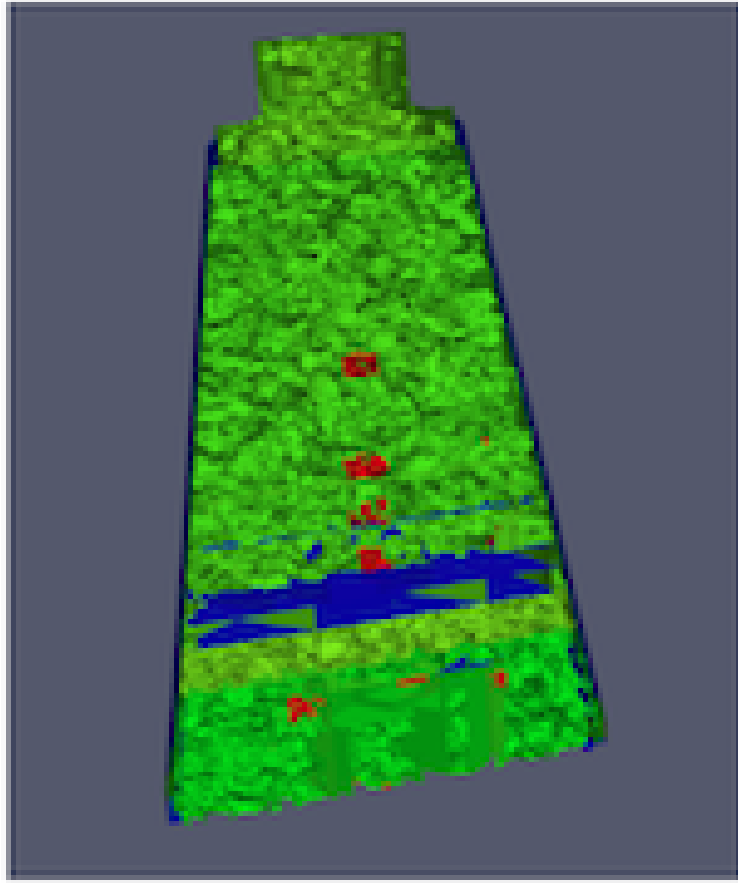
D3 Filter

The previous section described the importance of load balancing and ghost levels for parallel visualization. This section describes how to achieve that.

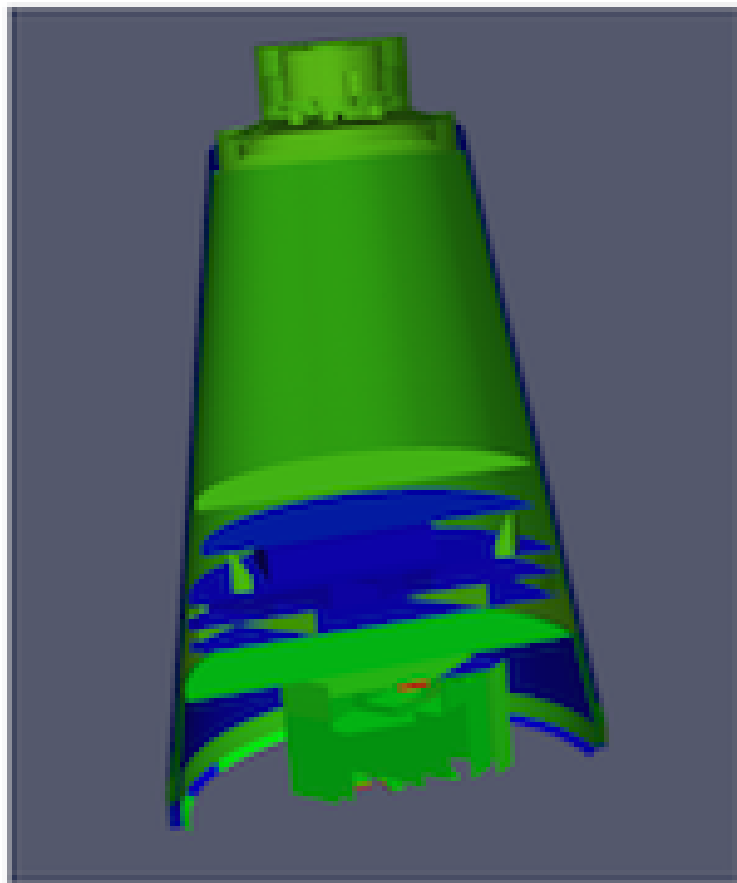
Load balancing and ghost cells are handled automatically by ParaView when you are reading structured data (image data, rectilinear grid, and structured grid). The implicit topology makes it easy to break the data into spatially coherent chunks and identify where neighboring cells are located.

It is an entirely different matter when you are reading in unstructured data (poly data and unstructured grid). There is no implicit topology and no neighborhood information available. ParaView is at the mercy of how the data was written to disk. Thus, when you read in unstructured data, there is no guarantee of how well-load balanced your data will be. It is also unlikely that the data will have ghost cells available, which means that the output of some filters may be incorrect.

Fortunately, ParaView has a filter that will both balance your unstructured data and create ghost cells. This filter is called D3, which is short for distributed data decomposition. Using D3 is easy; simply attach the filter (located in *Filters > Alphabetical > D3*) to whatever data you wish to repartition.



The most common use case for D3 is to attach it directly to your unstructured grid reader. Regardless of how well-load balanced the incoming data might be, it is important to be able to retrieve ghost cell so that subsequent filters will generate the correct data. The example above shows a cutaway of the extract surface filter on an unstructured grid. On the left, we see that there are many faces improperly extracted because we are missing ghost cells. On the right, the problem is fixed by first using the D3 filter.



2.6.7 Ghost Cells Generator

If your unstructured grid data is already partitioned satisfactorily but does not have ghost cells, it is possible to generate them using the **Ghost Cells Generator** filter. This filter can be attached to a source just like the D3 filter. Unlike D3, it will not repartition the dataset, it will only generate ghost cells, which is needed for some algorithms to execute correctly.

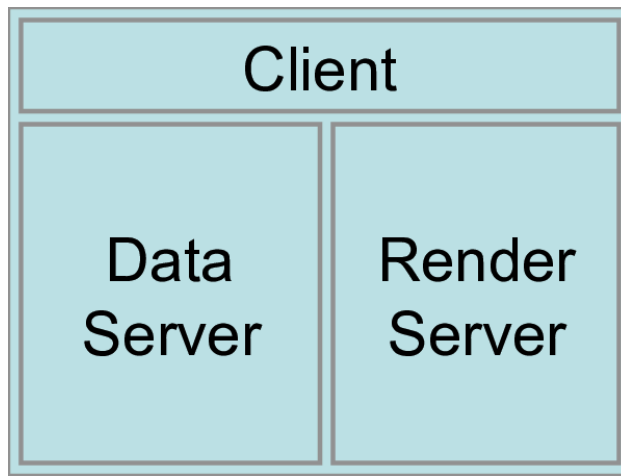
The **Ghost Cells Generator** has several options. **Build If Required** tells the filter to generate ghost cells only if required by a downstream filter. Since computing ghost cells is a computationally and communications intensive process, turning this option on can potentially save a lot of processing time. The **Minimum Number Of Ghost Levels** specifies at least how many ghost levels should be generated if **Build If Required** is off. Downstream filters may request more ghost levels than this minimum, in which case the **Ghost Cells Generator** will generate the requested number of ghost levels. The **Use Global Ids** option makes use of a **GlobalIds** array if it is present if on. If off, ghost cells are determined by coincident points.

2.6.8 ParaView architecture

Before we see how to use ParaView for parallel data processing, let's take a closer look at the ParaView architecture. ParaView is designed as a three-tier client-server architecture. The three logical units of ParaView are as follows.

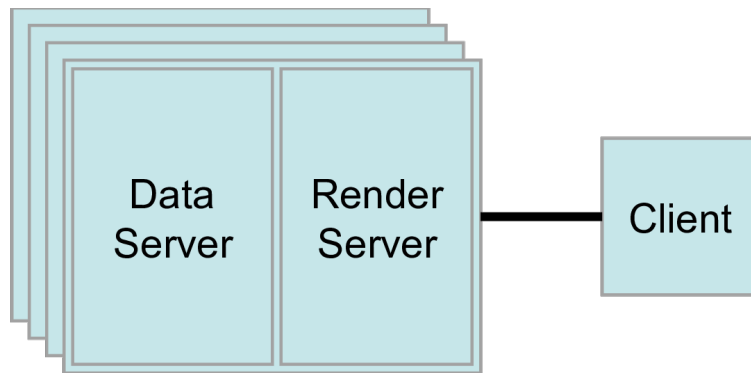
- **Data Server** The unit responsible for data reading, filtering, and writing. All of the pipeline objects seen in the pipeline browser are contained in the data server. The data server can be parallel.
- **Render Server** The unit responsible for rendering. The render server can also be parallel, in which case built-in parallel rendering is also enabled.
- **Client** The unit responsible for establishing visualization. The client controls the object creation, execution, and destruction in the servers, but does not contain any of the data (thus allowing the servers to scale without bottlenecking on the client). If there is a GUI, that is also in the client. The client is always a serial application.

These logical units need not be physically separated. Logical units are often embedded in the same application, removing the need for any communication between them. There are three modes in which you can run ParaView.

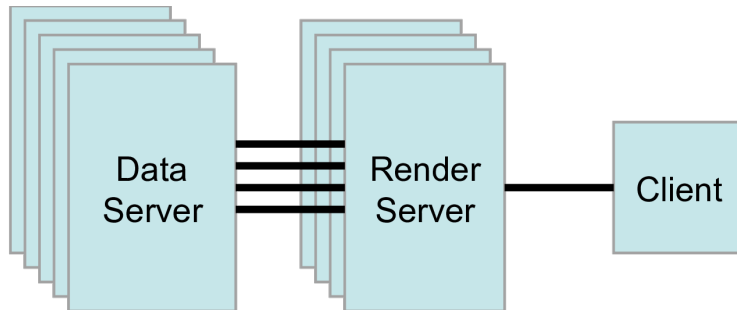


The first mode, with which you are already familiar, is **standalone** mode. In standalone mode, the client, data server, and render server are all combined into a single serial application. When you run the `paraview` application, you are automatically connected to a **builtin** server so that you are ready to use the full features of ParaView.

The second mode is **client-server** mode. In client-server mode, you execute the `pvserver` program on a parallel machine and connect to it with the `paraview` client application (or `pvc`). The `pvserver` program has both the data server and render server embedded in it, so both data processing and rendering take place there. The client and



server are connected via a socket, which is assumed to be a relatively slow mode of communication, so data transfer over this socket is minimized. We saw this mode of operation in [Section 2.6.2](#).



The third mode is **client-render server-data server** mode. In this mode, all three logical units are running in separate programs. As before, the client is connected to the render server via a single socket connection. The render server and data server are connected by many socket connections, one for each process in the render server. Data transfer over the sockets is minimized.

Although the client-render server-data server mode is supported, we almost never recommend using it. The original intention of this mode is to take advantage of heterogeneous environments where one might have a large, powerful computational platform and a second smaller parallel machine with graphics hardware in it. However, in practice, we find any benefit is almost always outstripped by the time it takes to move geometry from the data server to the render server. If the computational platform is much bigger than the graphics cluster, then use software rendering on the large computational platform. If the two platforms are about the same size, just perform all the computation on the graphics cluster. The executables used for this mode are `paraview` (or `pvpython`) (acting as the client), `pvdataserver` for the data-server, and `pvrenderserver` for the render-server.

2.6.9 Parallel processing in paraview and pvpython

To leverage parallel processing capabilities in `paraview` or `pvpython`, one has to use remote visualization, i.e., one has to connect to a `pvserver`. The processing for connecting to this `pvserver` is not different from what we say in [Section 2.6.2](#) and [Section 2.6.3](#). The only thing that changes is how the `pvserver` is launched.

You can start `pvserver` to run on more than one processing core using `mpirun` .

```
mpirun -np 4 pvserver
```

This will run `pvserver` on four processing cores. It will still listen for an incoming connection from a client on the default port. The big difference when running `pvserver` this way is that when data is loaded from a source, it will be distributed across the four cores if the data source is `parallel` aware and supports distributing the data across the different processing cores.

To see how this data is distributed, run `pvserver` as the command above and connect to it with `paraview`. Next, create another `Sphere` source using *Source > Sphere*. Change the array to color by to `vtkProcessId`. You will see an image like Figure Fig. 2.26.

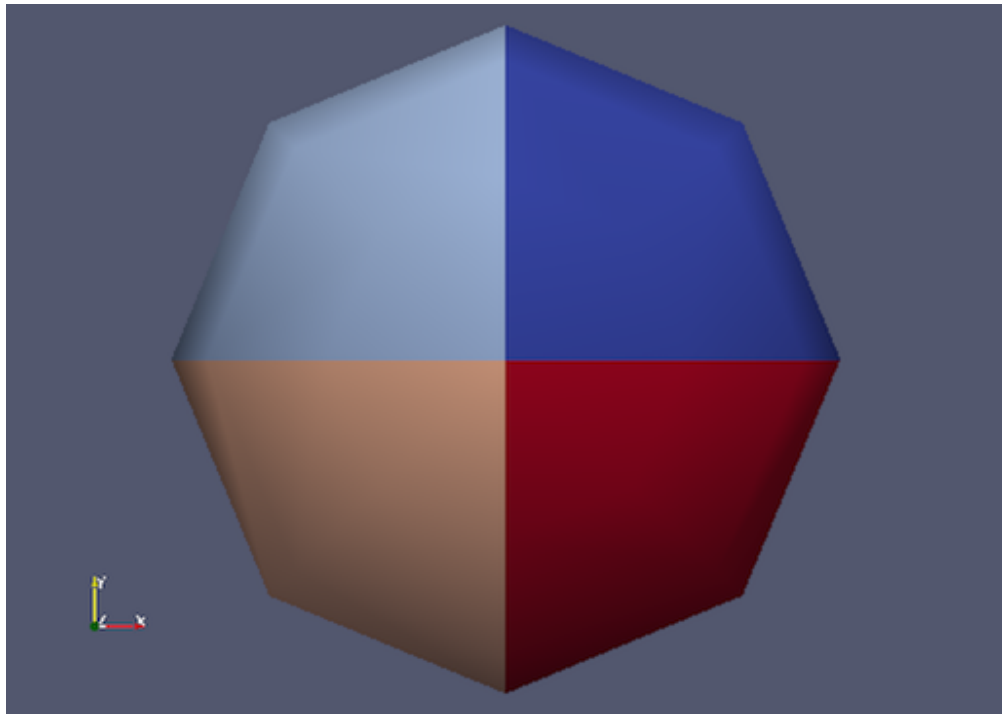


Fig. 2.26: Sphere source colored by `vtkProcessId` array that encodes the processing core on which the sphere data resides. Here, the sphere data is split among the four processing cores invoked by the command `mpirun -np 4 pvserver`.

If a data reader or source is not `parallel aware`, you can still get the benefits of spreading the data among processing cores by using the `D3` filter. This filter partitions a dataset into convex regions and transfers each region to a different processing core. To see an example of how `D3` partitions a dataset, create a *Source > Wavelet* while `paraview` is still connected to the `pvserver`. Next, select *Filters > Alphabetical > D3* and click `Apply`. The output of `D3` will not initially appear different from the original wavelet source. If you color by `vtkProcessId`, however, you will see the four partitions that have been distributed to the server processing cores.

2.6.10 Using `pbatch`

In Section 2.6.9, we said that to use parallel processing capabilities, one has to use remote visualization, i.e., one must use ParaView in a client-server mode with the client (`paraview` or `pvython`) connecting to a server (`pvserver`) that is being run in parallel using `mpirun`. However, there is one exception: `pbatch`. `pvython` and `pbatch` are quite similar in that both are similarly to the `python`

executable that can be used to run Python scripts. The extra thing that these executables do when compared with the standard `python` is that they initialize the environment so that any scripts that you run will be able to locate the ParaView Python modules and libraries automatically. `pvython` is exactly like the `paraview` executable without the GUI. You can think of it as the GUI from `paraview` is replaced by a Python interpreter in `pvython`. `pbatch`, on the other hand, can be thought of a `pvserver` where, instead of taking the control command from a remote client (`paraview` or `pvython`), in `pbatch`, the commands are taken from a Python script that is executed in the `pbatch` executable itself. Since `pbatch` is akin to the `pvserver`, unlike `pvython`, it can be run in parallel using `mpirun`. In that case, the root rank (or the first rank or the rank with index `0`) is the one that acts as the client, interpreting

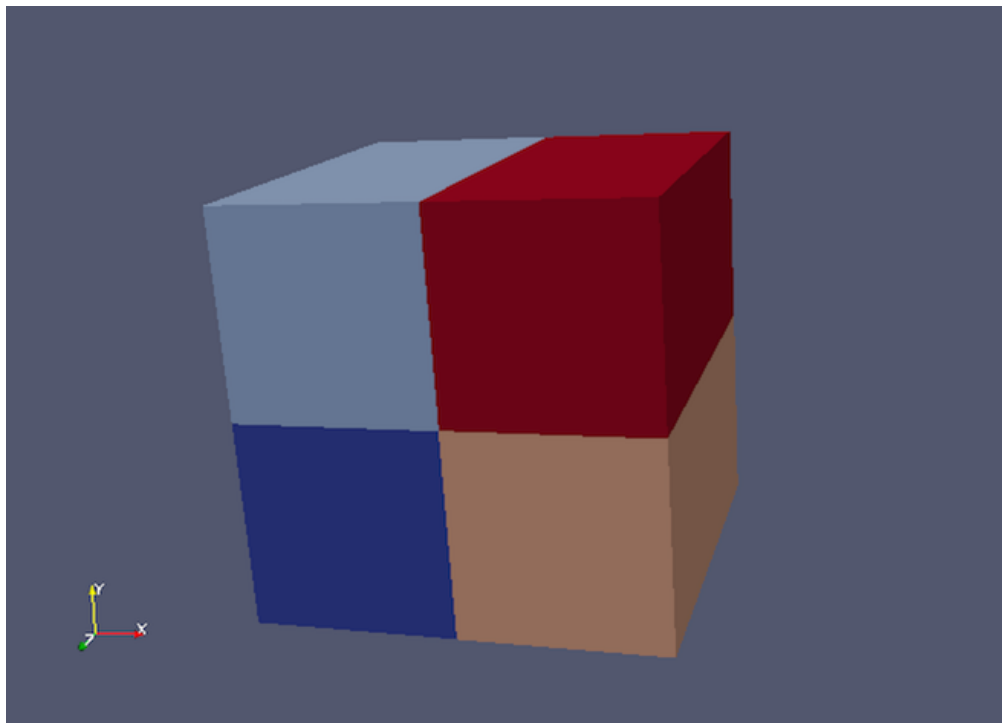


Fig. 2.27: Wavelet source processed by the D3 filter and colored by `vtkProcessId` array. Note how four regions of the image data are split evenly among the four processing cores when `pvserver` is run with `mpirun -np 4 pvserver`.

the Python script to execute the commands. Since `pbatch` is designed to act as its own server, you cannot connect to a remote server in the Python script, i.e., you cannot use `simple.Connect`. Furthermore, `pbatch` is designed for batch operation, which means that you can only specify the Python script as a command line argument. Unlike `pvython`, you cannot run this executable to get an interactive shell to enter Python commands.

```
# process the sample.py script in single process mode.
> pbatch sample.py

# process the sample.py script in parallel.
> mpirun -np 4 sample.py
```

In general, you should use `pvython` if you will be using the interpreter interactively and `pbatch` if you are running in parallel.

2.6.11 Fetching data to the client

Section 1.3.3 describes how to obtain information *about* a data object, but not how to access the data object itself. This section describes several ways to access data from within a Python script. The client/server nature of ParaView requires a couple steps to access the raw data. The Python script runs on the client side in either `pvython` or `paraview`, so one step involves moving the data from the server to the client. This can be accomplished with the following:

```
from paraview.simple import *
Connect("myhost")

# Create a sphere source on myhost
```

(continues on next page)

(continued from previous page)

```
s = Sphere()
full_sphere = servermanager.Fetch(s)
```

Here, the full dataset is moved from the server to the client.

The second step is required to deal with the fact that data on the remote server may be split across distributed processes. By default, `servermanager.Fetch(s)` appends all the pieces on the different remote processes and produces the appended data set on the client. The exact append operation depends on the type of dataset being retrieved. Composite datasets are merged by treating the dataset piece on each distributed process as a block merged into a new multiblock dataset, polygonal datasets are appended into a single polygonal dataset, rectilinear grids are appended into a single rectilinear grid, and other datasets are appended into an unstructured grid. Distributed image datasets cannot currently be fetched to the client. Care must be taken when fetching an entire dataset to the client because the data that fits on many distributed processes on a remote system may not fit in client memory.

Another option is to fetch just a single piece of the dataset on one remote process to the client. To do this, pass the rank number of the remote process from which you want to retrieve the data to the `Fetch` function, e.g.,

```
# Retrieve the piece of the data set on remote process 2
s = Sphere()
sphere_piece = servermanager.Fetch(s, 2)
```

Lastly, `servermanager.Fetch` provides a way to apply helper filters to the dataset that run at two stages. The filter for the first stage is applied to the data on each remote process, and the filter for the second stage is applied to the results from the first stage after they are gathered to the root server process. The results from the second stage of filtering are then transferred from the root server process to the client.

In the next example, the `Extract Surface` filter is applied to a source with data on each process in the first stage. The results are then assembled with the `Append Geometry` filter and sent to the client.

```
s = Sphere()
extract = servermanager.filters.ExtractSurface()
append = servermanager.filters.AppendGeometry()
full_surface = servermanager.Fetch(s, extract, append)
```

The second filter must be able to accept multiple connections and handle the output dataset type from the first filter.

2.6.12 Rendering

Rendering is the process of synthesizing the images that you see based on your data. The ability to effectively interact with your data depends highly on the speed of the rendering. Thanks to advances in 3D hardware acceleration, fueled by the computer gaming market, we have the ability to render 3D quickly even on moderately-priced computers. But, of course, the speed of rendering is proportional to the amount of data being rendered. As data gets bigger, the rendering process naturally gets slower.

To ensure that your visualization session remains interactive, ParaView supports two modes of rendering that are automatically flipped as necessary. In the first mode, **still render**, the data is rendered at the highest level of detail. This rendering mode ensures that all of the data is represented accurately. In the second mode, **interactive render**, speed takes precedence over accuracy. This rendering mode endeavors to provide a quick rendering rate regardless of data size.

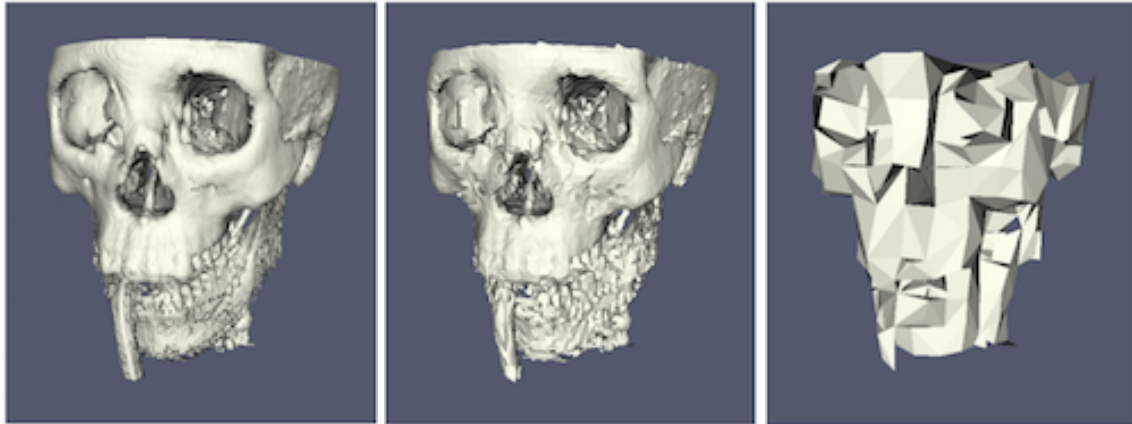
While you are interacting with a 3D view (for example, rotating, panning, or zooming with the mouse), ParaView uses an interactive render. This is because, during the interaction, a high frame rate is necessary to make these features usable and because each frame is immediately replaced with a new rendering while the interaction is occurring so that fine details are less important during this mode. At any time when interaction of the 3D view is not taking place, ParaView uses a still render so that the full detail of the data is available as you study it. As you drag your mouse in

a 3D view to move the data, you may see an approximate rendering. The full detail will be presented as soon as you release the mouse button.

The interactive render is a compromise between speed and accuracy. As such, many of the rendering parameters concern when and how lower levels of detail are used.

Basic Rendering Settings

Some of the most important rendering options are the LOD parameters. During interactive rendering, the geometry may be replaced with a lower **level of detail (LOD)**, an approximate geometry with fewer polygons.



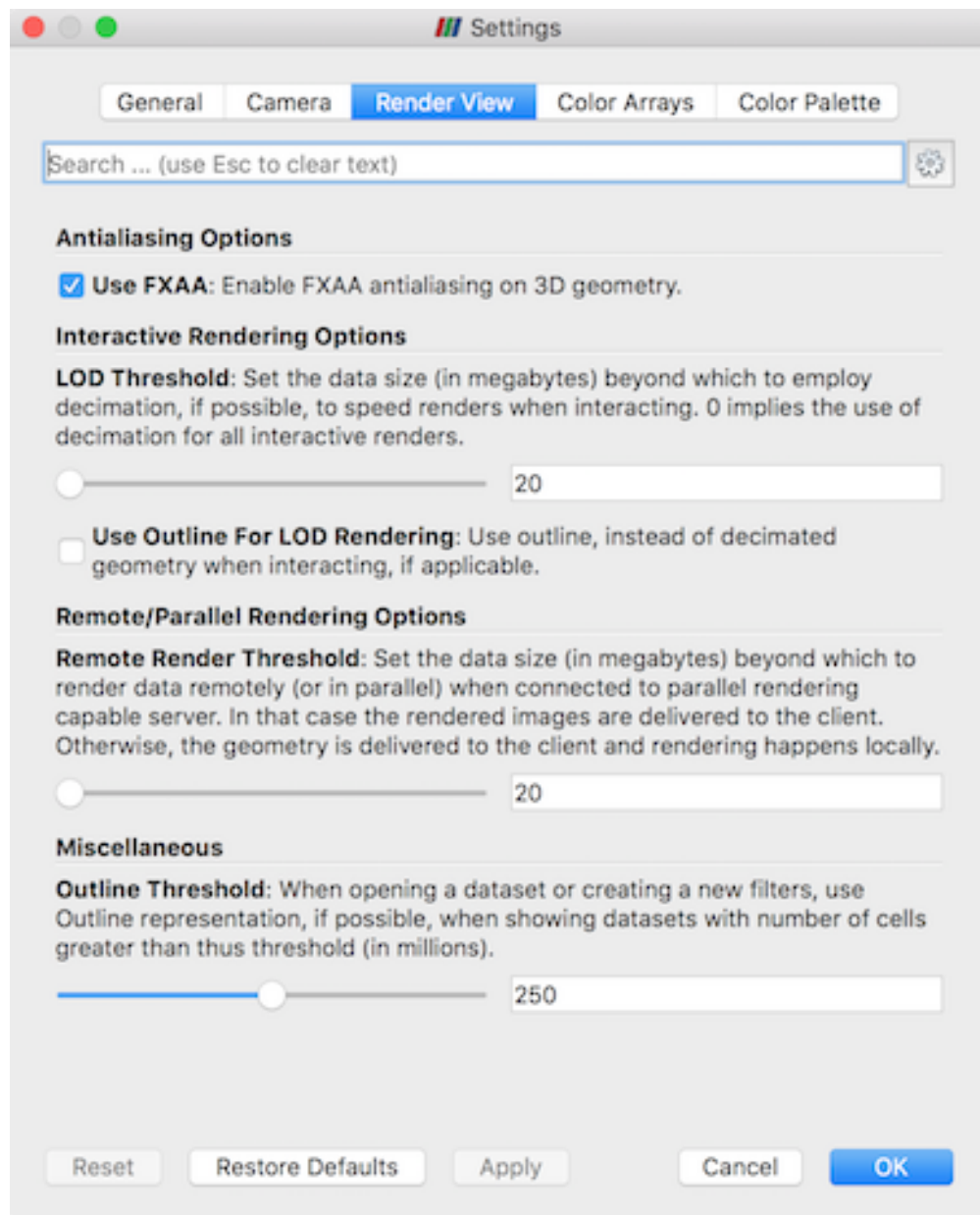
The resolution of the geometric approximation can be controlled. In the proceeding images, the left image is the full resolution, the middle image is the default decimation for interactive rendering, and the right image is ParaView's maximum decimation setting.

The 3D rendering parameters are located in the settings dialog box, which is accessed in the menu from the *Edit > Settings* menu (*ParaView > Preferences* on the Mac). The rendering options in the dialog are in the **Render View** tab.

The options pertaining to the geometric decimation for interactive rendering are located in a section labeled **Interactive Rendering Options**. Some of these options are considered advanced, so to access them, you have to either toggle on the advanced options with the button or search for the option using the edit box at the top of the dialog. The interactive rendering options include the following.

- **LOD Threshold** : Set the data size at which to use a decimated geometry in interactive rendering. If the geometry size is under this threshold, ParaView always renders the full geometry. Increase this value if you have a decent graphics card that can handle larger data. Try decreasing this value if your interactive renders are too slow.
- **LOD Resolution** : Set the factor that controls how large the decimated geometry should be. This control is set to a value between 0 and 1. 0 produces a very small number of triangles but, possibly, with a lot of distortion. 1 produces more detailed surfaces but with larger geometry.
- **Non Interactive Render Delay** : Add a delay between an interactive render and a still render. ParaView usually performs a still render immediately after an interactive motion is finished (for example, releasing the mouse button after a rotation). This option can add a delay that can give you time to start a second interaction before the still render starts, which is helpful if the still render takes a long time to complete.
- **Use Outline For LOD Rendering** : Use an outline in place of decimated geometry. The outline is an alternative for when the geometry decimation takes too long or still produces too much geometry. However, it is more difficult to interact with just an outline.

ParaView contains many more rendering settings. Here is a summary of some other settings that can effect the rendering performance regardless of whether ParaView is run in client-server mode or not. These options are spread among



several categories, and several are considered advanced.

- **Translucent Rendering Options**

- Depth Peeling : Enable or disable depth peeling. Depth peeling is a technique ParaView uses to properly render translucent surfaces. With it, the top surface is rendered and then “peeled away” so that the next lower surface can be rendered and so on. If you find that making surfaces transparent really slows things down or renders completely incorrectly, then your graphics hardware may not be implementing the depth peeling extensions well; try shutting off depth peeling. 
- Depth Peeling for Volumes : Include volumes in depth peeling to correctly intermix volumes and translucent polygons.
- Maximum Number Of Peels : Set the maximum number of peels to use with depth peeling. Using more peels allows more depth complexity, but allowing less peels runs faster. You can try adjusting this parameter if translucent geometry renders too slow or translucent images do not look correct. 

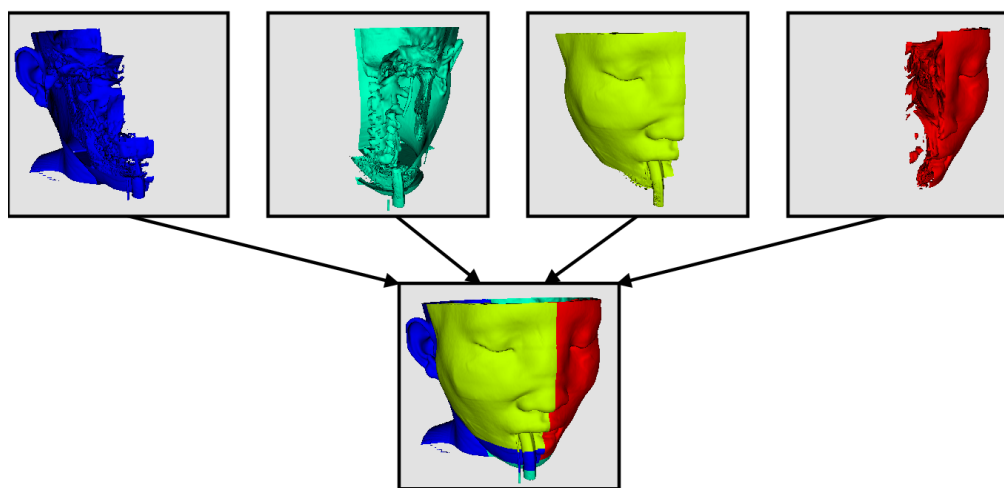
- **Miscellaneous**

- Outline Threshold : When creating very large datasets, default to the outline representation. Surface representations usually require ParaView to extract geometry of the surface, which takes time and memory. For data with sizes above this threshold, use the outline representation, which has very little overhead, by default instead.
- Show Annotation : Show or hide annotation providing rendering performance information. This information is handy when diagnosing performance problems. 

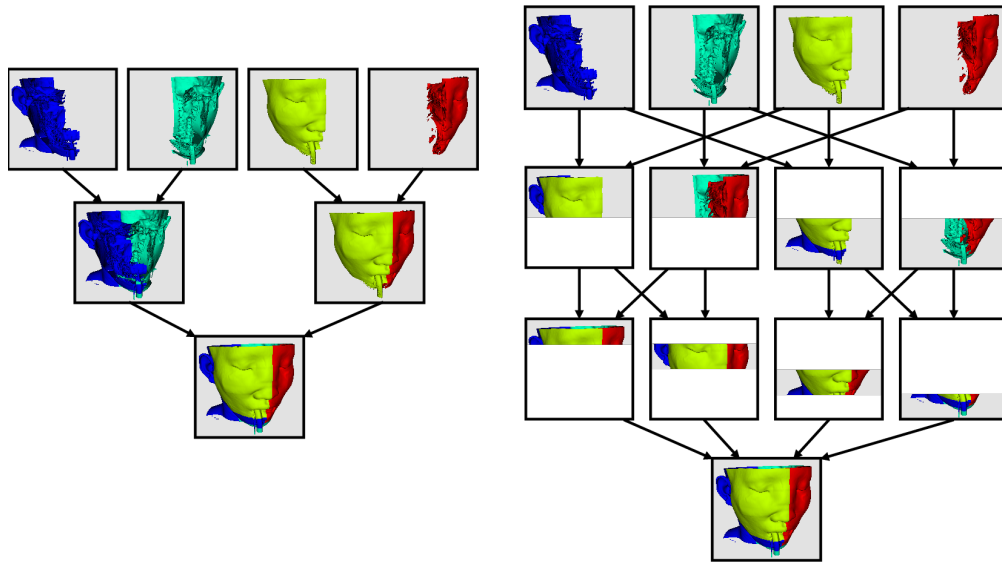
Note that this is not a complete list of ParaView rendering settings. We have left out settings that do not significantly affect rendering performance. We have also left out settings that are only valid for parallel client-server rendering, which are discussed in [Section 2.6.12](#).

Basic Parallel Rendering

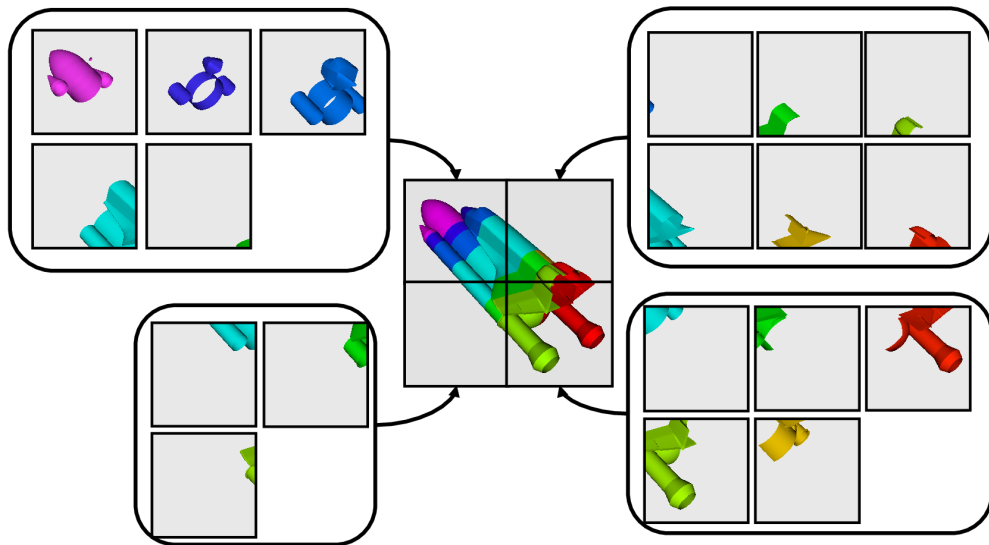
When performing parallel visualization, we are careful to ensure that the data remains partitioned among all of the processes up to and including the rendering processes. ParaView uses a parallel rendering library called **IceT**. IceT uses a **sort-last** algorithm for parallel rendering. This parallel rendering algorithm has each process independently render its partition of the geometry and then **composites** the partial images together to form the final image.



The preceding diagram is an oversimplification. IceT contains multiple parallel image compositing algorithms such as **binary tree**, **binary swap**, and **radix-k** that efficiently divide work among processes using multiple phases.



The wonderful thing about sort-last parallel rendering is that its efficiency is completely insensitive to the amount of data being rendered. This makes it a very scalable algorithm and well suited to large data. However, the parallel rendering overhead does increase linearly with the number of pixels in the image. Consequently, some of the rendering parameters deal with the image size.



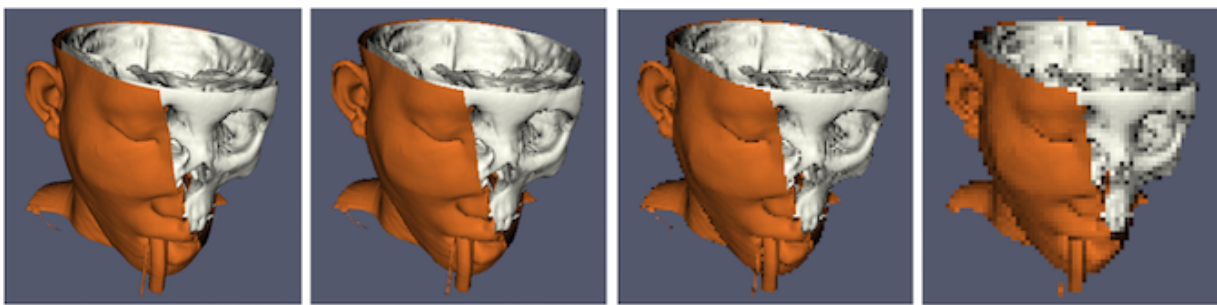
IceT also has the ability to drive tiled displays, which are large, high-resolution displays comprising an array of monitors or projectors. Using a sort-last algorithm on a tiled display is a bit counterintuitive because the number of pixels to composite is so large. However, IceT is designed to take advantage of spatial locality in the data on each process to drastically reduce the amount of compositing necessary. This spatial locality can be enforced by applying the *Filters > Alphabetical > D3* filter to your data.

Because there is an overhead associated with parallel rendering, ParaView has the ability to turn off parallel rendering at any time. When parallel rendering is turned off, the geometry is shipped to the location where display occurs. Obviously, this should only happen when the data being rendered is small.

Image Level of Detail

The overhead incurred by the parallel rendering algorithms is proportional to the size of the images being generated. Also, images generated on a server must be transferred to the client, a cost that is also proportional to the image size. To help increase the frame rate during interaction, ParaView introduces a new LOD parameter that controls the size of the images.

During interaction while parallel rendering, ParaView can optionally **subsample** the image. That is, ParaView will reduce the resolution of the image in each dimension by a factor during interaction. Reduced images will be rendered, composited, and transferred. On the client, the image is inflated to the size of the available space in the GUI.



The resolution of the reduced images is controlled by the factor with which the dimensions are divided. In the proceeding images, the left image has the full resolution. The following images were rendered with the resolution reduced by a factor of 2, 4, and 8, respectively.

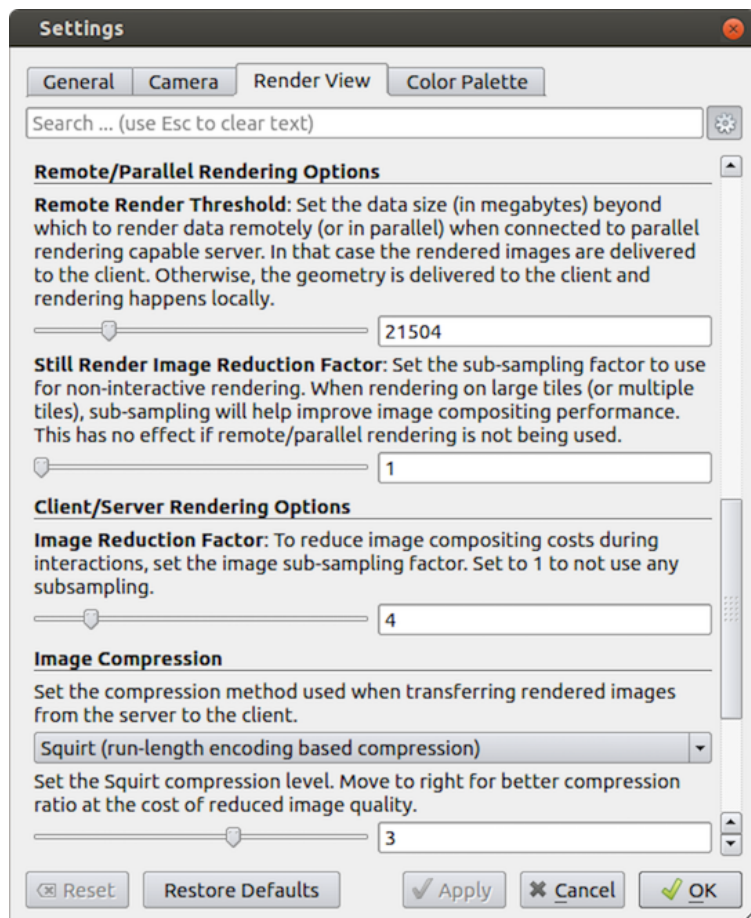
ParaView also has the ability to compress images before transferring them from server to client. Compression, of course, reduces the amount of data transferred and, therefore, makes the most of the available bandwidth. However, the time it takes to compress and decompress the images adds to the latency.

ParaView contains several different image compression algorithms for client-server rendering. The first uses LZ4 compression that is designed for high-speed compression and decompression. The second option is a custom algorithm called **Squirt**, which stands for Sequential Unified Image Run Transfer. Squirt is a run-length encoding compression that reduces color depth to increase run lengths. The third algorithm uses the **Zlib** compression library, which implements a variation of the Lempel-Ziv algorithm. Zlib typically provides better compression than Squirt, but it takes longer to perform and, hence, adds to the latency. paraview Windows and Linux executables include a compression option that uses NVIDIA's NVPipe library for hardware-accelerated compression and decompression if a Kepler-class or higher NVIDIA GPU is available.

Parallel Render Parameters

Like the other 3D rendering parameters, the parallel rendering parameters are located in the **Settings** dialog. The parallel rendering options in the dialog are in the **Render View** tab (intermixed with several other rendering options such as those described in [Section 2.6.12](#)). The parallel and client-server options are divided among several categories, and several are considered advanced.

- Remote/Parallel Rendering Options
 - **Remote Render Threshold**: Set the data size at which to render remotely in parallel or to render locally. If the geometry is over this threshold (and ParaView is connected to a remote server), the data is rendered in parallel remotely, and images are sent back to the client. If the geometry is under this threshold, the geometry is sent back to the client, and images are rendered locally on the client.
 - **Still Render Image Reduction Factor**: Set the sub-sampling factor for still (non-interactive) rendering. Some large displays have more resolution than is really necessary, so this sub-sampling reduces the resolution of all images displayed. 
- Client/Server Rendering Options



- **Image Reduction Factor**: Set the interactive subsampling factor. The overhead of parallel rendering is proportional to the size of the images generated. Thus, you can speed up interactive rendering by specifying an image subsampling rate. When this box is checked, interactive renders will create smaller images, which are then magnified when displayed. This parameter is only used during interactive renders. 
- **Image Compression**
 - Before images are shipped from server to client, they can optionally be compressed using one of three available compression algorithms: LZ4 , Squirt , or Zlib . To make the compression more effective, either algorithm can reduce the color resolution of the image before compression. The sliders determine the amount of color bits saved. Full color resolution is always used during a still render. 
 - Suggested image compression presets are provided for several common network types. When attempting to select the best image compression options, try starting with the presets that best match your connection. 

Parameters for Large Data

The default rendering parameters are suitable for most users. However, when dealing with very large data, it can help to tweak the rendering parameters. While the optimal parameters depend on your data and the hardware on which ParaView is running, here are several pieces of advice that you should follow.

- If there is a long pause before the first interactive render of a particular dataset, it might be the creation of the decimated geometry. Try using an outline instead of decimated geometry for interaction. You could also try lowering the factor of the decimation to 0 to create smaller geometry.
- Avoid shipping large geometry back to the client. The remote rendering will use the power of the entire server to render and ship images to the client. If remote rendering is off, geometry is shipped back to the client. When you have large data, it is always faster to ship images than to ship data. (Although, if your network has a high latency, this could become problematic for interactive frame rates.)
- Adjust the interactive image sub-sampling for client-server rendering as needed. If image compositing is slow, if the connection between client and server has low bandwidth, or if you are rendering very large images, then a higher subsample rate can greatly improve your interactive rendering performance.
- Make sure **Image Compression** is on. It has a tremendous effect on desktop delivery performance, and the artifacts it introduces, which are only there during interactive rendering, are minimal. Lower bandwidth connections can try using Zlib instead of Squirt compression. Zlib will create smaller images at the cost of longer compression/decompression times.
- If the network connection has a high latency, adjust the parameters to avoid remote rendering during interaction. In this case, you can try turning up the remote rendering threshold a bit, and this is a place where using the outline for interactive rendering is effective.
- If the still (non-interactive) render is slow, try turning on the delay between interactive and still rendering to avoid unnecessary renders.

2.7 Memory Inspector

The **ParaView** Memory Inspector panel provides users with a convenient way to monitor **ParaView**'s memory usage during interactive visualization. It also provides developers with a point-and-click interface for attaching a debugger to local or remote client and server processes. As explained earlier, both the Information panel and the Statistics Inspector are prone to over and under estimate the total memory used for the current pipeline. The Memory Inspector addresses those issues through direct queries to the operating system. A number of diagnostic statistics are gathered and reported, including the total memory used by all processes on a per-host basis, the total cumulative memory use by **ParaView** on a per-host basis, and the individual per-rank use by each **ParaView** process. When memory consumption reaches a critical level, either cumulatively on the host or in an individual rank, the corresponding GUI element will turn red, alerting you that you are in danger of potentially being shut down. This gives you a chance to save state and restart the job with more nodes to avoid losing your work. On the flip side, knowing when you're not close to using the full capacity of available memory can be useful to conserve computational resources by running smaller jobs. Of course, the memory foot print is only one factor in determining the optimal run size.

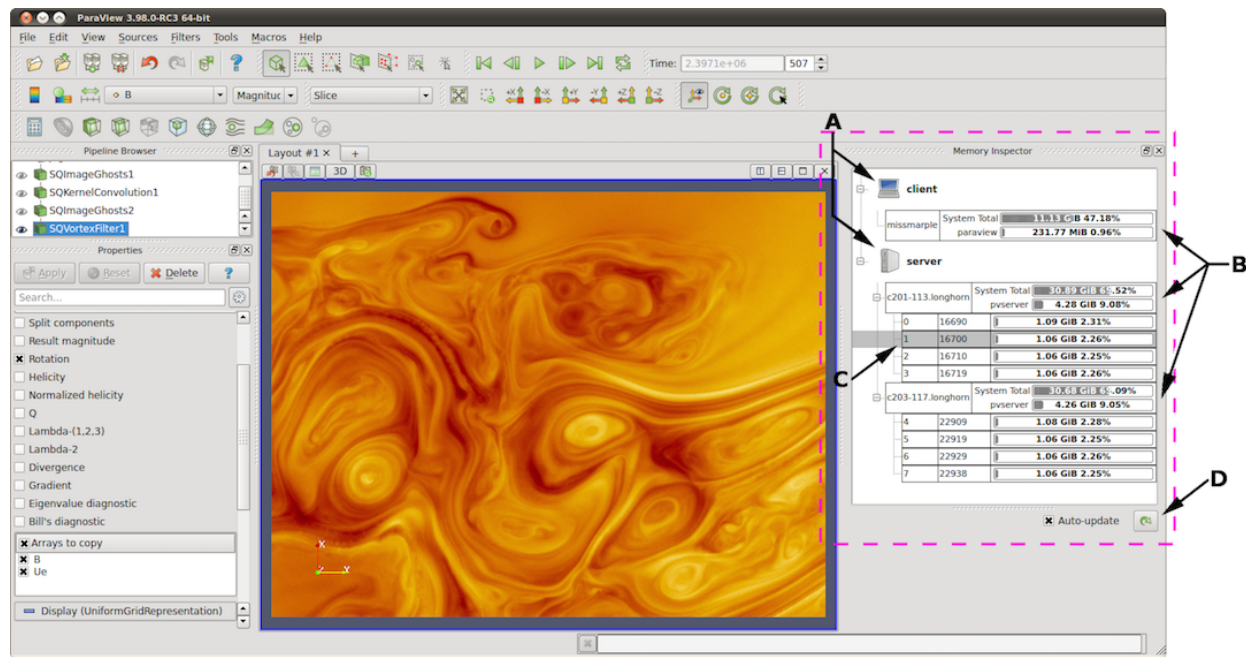


Fig. 2.28: The main UI elements of the Memory Inspector panel. A: Process Groups, B: Per-Host statistics, C: Per-Rank statistics, and D: Update controls.

2.7.1 User interface and layout

The Memory Inspector panel displays information about the current memory usage on the client and server hosts. Fig. 2.28 shows the main UI elements labeled A-D. A number of additional features are provided via specialized context menus accessible from the Client and Server group, Host, and Rank's UI elements. The main UI elements are:

A. Process Groups

1. *Client* : There is always a client group that reports statistics about the **ParaView** client.
2. *Server* : When running in client-server mode, a server group reports statistics about the hosts where **pvserver** processes are running.
3. *Data Server* : When running in client-data-render server mode, a data server group reports statistics about the hosts where **pvdataserver** processes are running.

4. *Render Server* : When running in client-data-render server mode, a render server group reports statistics about the hosts where `pvrenderserver` processes are running.
- B. *Per-Host Statistics* : Per-host statics are reported for each host where a **ParaView** process is running. Hosts are organized by host name, which is shown in the first column. Two statics are reported: 1) total memory used by all processes on the host and 2) **ParaView**'s cumulative usage on this host. The absolute value is printed in a bar that shows the percentage of the total available memory used. On systems where job-wide resource limits are enforced, **ParaView** is made aware of the limits via the `PV_HOST_MEMORY_LIMIT` environment variable, in which case, **ParaView**'s cumulative percent used is computed using the smaller of the host total and the resource limit.
- C. *Update Controls* : By default, when the panel is visible, memory use statistics are updated automatically as pipeline objects are created, modified, or destroyed and after the scene is rendered. Updates may be triggered manually by using the refresh button. Automatic updates may be disabled by un-checking the *Auto-update* check box. Queries to remote systems have proven to be very fast even for fairly large jobs. Hence, the auto-update feature is enabled by default.
- D. *Host Properties Dialog* : The *Host* context menu provides the *Host Properties* dialog, which reports various system details such as the OS version and the CPU version, as well as the memory installed and available to the host context and process context. While the *Memory Inspector* panel reports memory use as a percent of the available in the given context, the *Host Properties* dialog reports the total memory installed and available in each context. Comparing the installed and available memory can be used to determine if you are impacted by resource limits.

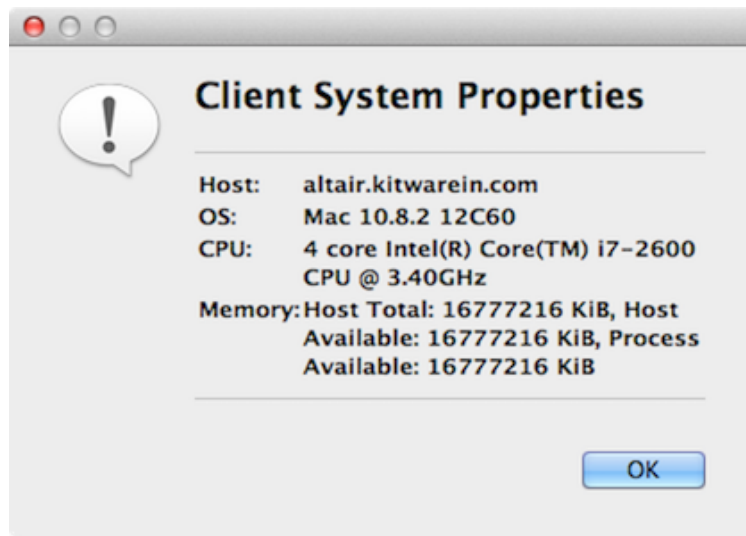


Fig. 2.29: Host properties dialog.

2.7.2 Advanced debugging features

Remote commands

The *Memory Inspector Panel* provides a remote (or local) command feature, allowing you to execute a shell command on a given host. This feature is exposed via a specialized Rank item context menu. Because we have information such as a rank's process id, individual processes may be targeted. For example, this allows you to quickly attach a debugger to a server process running on a remote cluster. If the target rank is not on the same host as the client, then the command is considered remote. Otherwise, it is considered local. Therefore, remote commands are executed via `ssh`, while local commands are not. A list of command templates is maintained. In addition to a number of pre-defined command templates, you may add templates or edit existing ones. The default templates allow you to:

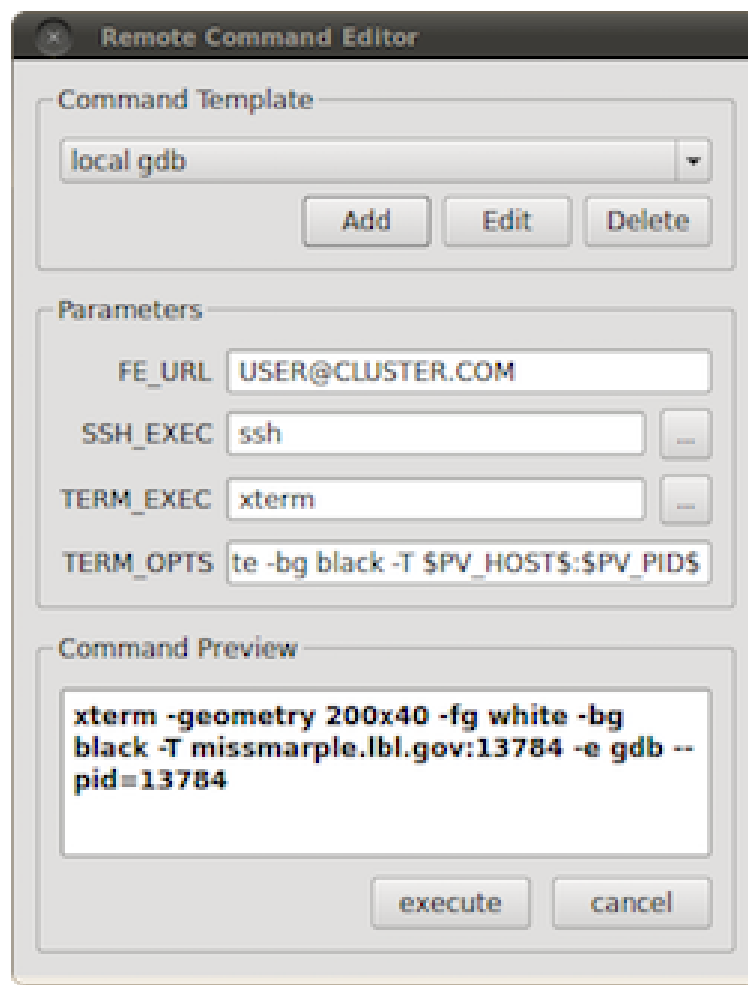


Fig. 2.30: The remote command dialog.

- Attach gdb to the selected process
- Run top on the host of the selected process
- Send a signal to the selected process

Prior to execution, the selected template is parsed, and a list of special tokens are replaced with runtime-determined or user-provide values. User-provided values can be set and modified in the dialog's parameter group. The command, with tokens replaced, is shown for verification in the dialog's preview pane.

The following tokens are available and may be used in command templates as needed:

1. `$TERM_EXEC$` : The terminal program that will be used to execute commands. On Unix systems, xterm is typically used. On Windows systems, `cmd.exe` is typically used. If the program is not in the default path, then the full path must be specified.
2. `$TERM_OPTS$` : Command line arguments for the terminal program. On Unix, these may be used to set the terminals window title, size, colors, and so on.
3. `SSH_EXEC` : The program to use to execute remote commands. On Unix, this is typically ssh. On Windows, one option is `plink.exe`. If the program is not in the default path, then the full path must be specified.
4. `FE_URL` : Ssh URL to use when the remote processes are on compute nodes that are not visible to the outside world. This token is used to construct command templates where two ssh hops are made to execute the command.
5. `PV_HOST` : The hostname where the selected process is running.
6. `PV_PID` : The process-id of the selected process.

Note: On Windows, the debugging tools found in Microsoft's SDK need to be installed in addition to Visual Studio (e.g., `windbg.exe`). The ssh program `plink.exe` for Windows doesn't parse ANSI escape codes that are used by Unix shell programs. In general, the Windows- specific templates need some polishing.

Stack trace signal handler

The Process Group's context menu provides a back trace signal handler option. When enabled, a signal handler is installed that will catch signals such as SEGV, TERM, INT, and ABORT and that will print a stack trace before the process exits. Once the signal handler is enabled, you may trigger a stack trace by explicitly sending a signal. The stack trace signal handler can be used to collect information about crashes or to trigger a stack trace during deadlocks when it's not possible to ssh into compute nodes. Sites that restrict users' ssh access to compute nodes often provide a way to signal running processes from the login node. Note that this feature is only available on systems that provide support for POSIX signals, and we currently only have implemented stack trace for GNU-compatible compilers.

2.7.3 Compilation and installation considerations

If the system on which **ParaView** will run has special resource limits enforced, such as job-wide memory use limits, or non-standard per-process memory limits, then the system administrators need to provide this information to the running instances of **ParaView** via the following environment variables. For example, those could be set in the batch system launch scripts.

1. `PV_HOST_MEMORY_LIMIT` : For reporting host-wide resource limits.
2. `PV_PROC_MEMORY_LIMIT` : For reporting per-process memory limits. that are not enforced via standard Unix resource limits.

A few of the debugging features (such as printing a stack trace) require debug symbols. These features will work best when **ParaView** is built with `CMAKE_BUILD_TYPE=Debug` or, for release builds, `CMAKE_BUILD_TYPE=RelWithDebugSymbols`.

2.8 Multiblock Inspector

Composite datasets (Section 1.3) such as multiblock datasets and AMR are often encountered when visualizing results from several scientific simulation codes e.g. OpenFOAM, Exodus, etc. Readers for several of these simulation file formats support selecting which blocks to read. Additionally, you may want to control display properties such as visibility, opacity, and color for individual blocks or subtress. You can use the **Multiblock Inspector** panel for this.

Fig. 2.31 shows the **Multiblock Inspector** showing the hierarchy from an Exodus dataset. The panel tracks the active source and reflects the structure for the data produced by the active source. The display properties reflect their state in the active view.

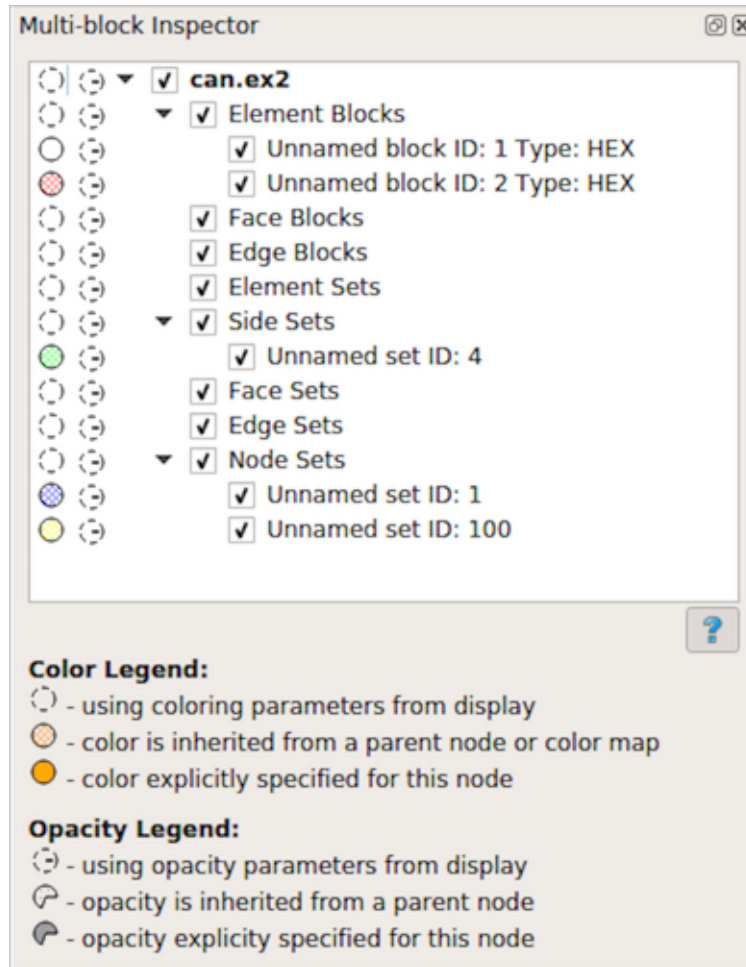


Fig. 2.31: The Multiblock Inspector

To show or hide a block you have to click on the checkbox next to the block name. Toggling the visibility of a non-leaf block, affects the visibility of the entire subtree.

The first column reflects the color used for the block. The color for a block can be specified in multiple ways. First, you can choose to not use any block specific overrides and simply let the default display properties (Section 1.4.3) affect the rendering. This is the default behavior. When that is the case for any block, it is indicated in the color column by an empty dotted circle . Second, you can explicitly override the color to use for a block or a subtree. To do this, simply double click on the color column next to the block of interest. This will pop up the color chooser dialog. Explicitly overridden color for a block is indicated by a solid filled circle icon filled with the selected color. When an explicit

color is set on a non-leaf node, all its children (and their children) inherit that color unless explicitly overridden. For such nodes that have a color inherited from their parent, we use the dotted-circle icon filled with a pattern icon .

The second column reflects the opacity override for the blocks. Similar to color, the opacity could be simply using value from the display properties , or explicitly set , or inherited from parent node .

Did you know?

In most cases with multiblock datasets, ParaView uses the `vtkBlockColors` array for coloring. This is an array filled with random values so that each block can be colored using a different color. That makes it easier to visually see each of the blocks in the view. When in this mode, the **Multiblock Inspector** ‘s color column shows the color used for each of the blocks using the same icon as the one used for inherited colors i.e. .

You can change colors and opacities for a specific block by double clicking on the corresponding icon. This allows setting values one at a time. For specifying color and opacity overrides for multiple elements, you can select the items and then right click to get the context menu. The context menu allows you to change these properties for all the selected items, as shown in Fig. 2.32.

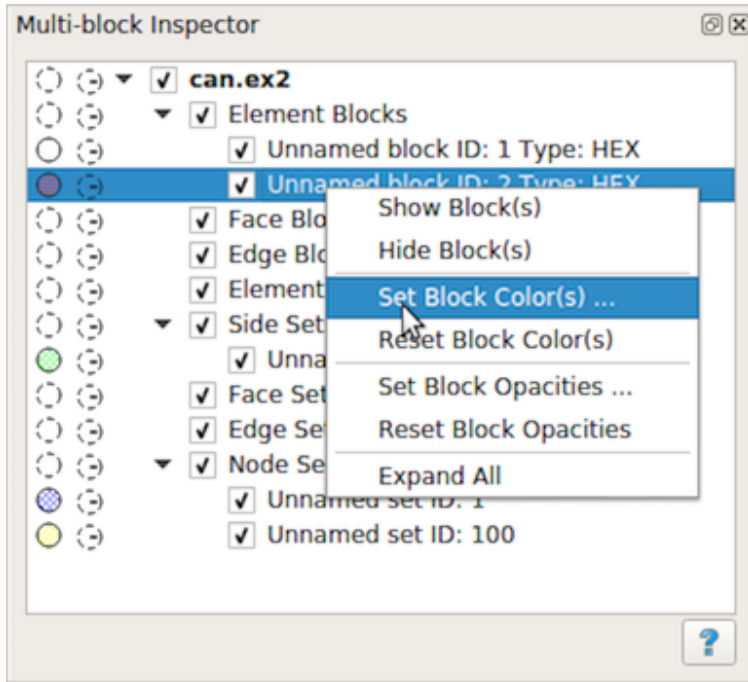


Fig. 2.32: The **Multiblock Inspector** context menu.

Besides using the **Multiblock Inspector** to set color and opacity overrides for blocks, you can also directly changes these parameters from the **Render View** itself. Simply right-click in the render view on the block of interest and you’ll get a context menu, Fig. 2.33, that allows changing the block properties and are more.

Another useful feature with the **Multiblock Inspector** is selection. If you simply click on any row in the inspector, you will select the block (or subtree) and the active view will highlight the selected block(s). Conversely, if you make a block-based selection in the active **Render View** using , you will see the corresponding blocks highlighted in the **Multiblock Inspector** panel.

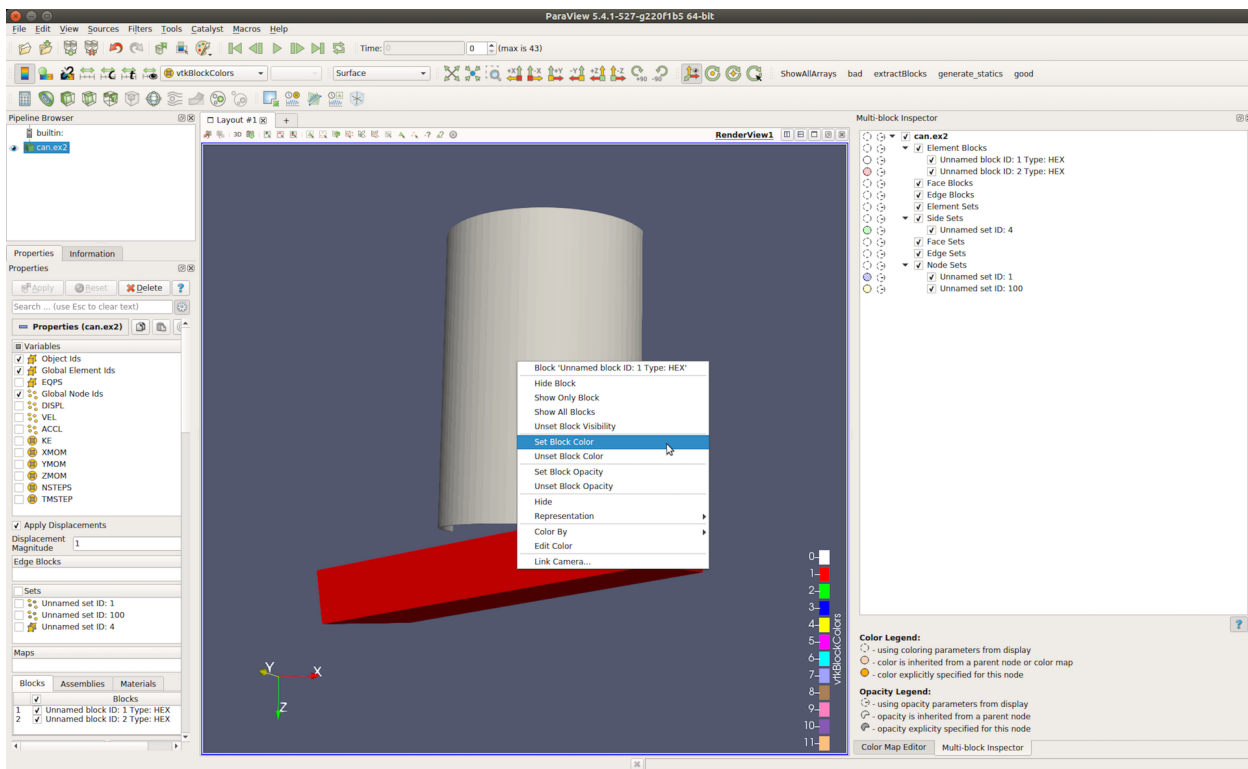


Fig. 2.33: Context menu in Render View can be used to change block display properties.

2.9 Annotations

Explicit labeling and annotation of particular data values is often an important element in data visualization and analysis. ParaView provides a variety of mechanisms to enable annotation in renderings ranging from free floating text rendered alongside other visual elements in the render view to data values associated with particular points or cells.

2.9.1 Annotation sources

Several types of text annotations can be added through the *Sources > Alphabetical* menu. Text from these sources is drawn on top of 3D elements in the render view. All annotation sources share some common properties under the **Display** section of the **Properties** panel. These include **Font Properties** such as the font to use, the size of the text, its color, opacity, and justification, as well as text effects to apply such as making it bold, italic, or shadowed.



Fig. 2.34: Font property controls in annotation sources and filters.

There are three fonts available in **ParaView**: Arial, Courier, and Times. You can also supply an arbitrary TrueType font file (*.ttf) to use by selecting the **File** entry in the popup menu under **Font Properties** and clicking on the **...** button to the right of the font file text field. A file selection dialog will appear letting you choose a font file from the file system on which `paraview` (or `pvc`) is running.

The remaining display properties control where the text is placed in the render view. There are two modes for placement,

one that uses predefined positions relative to the render view, and one that enables arbitrary interactive placement in the render view. The first mode is active when the `Use Window Location` checkbox is selected. It enables the annotation to be placed in one of the four corners of the render view or centered horizontally at the top or bottom of the render view. Buttons with icons representing the location are shown in the Pipeline browser. These buttons correspond to locations in the render view as depicted in Fig. 2.35.

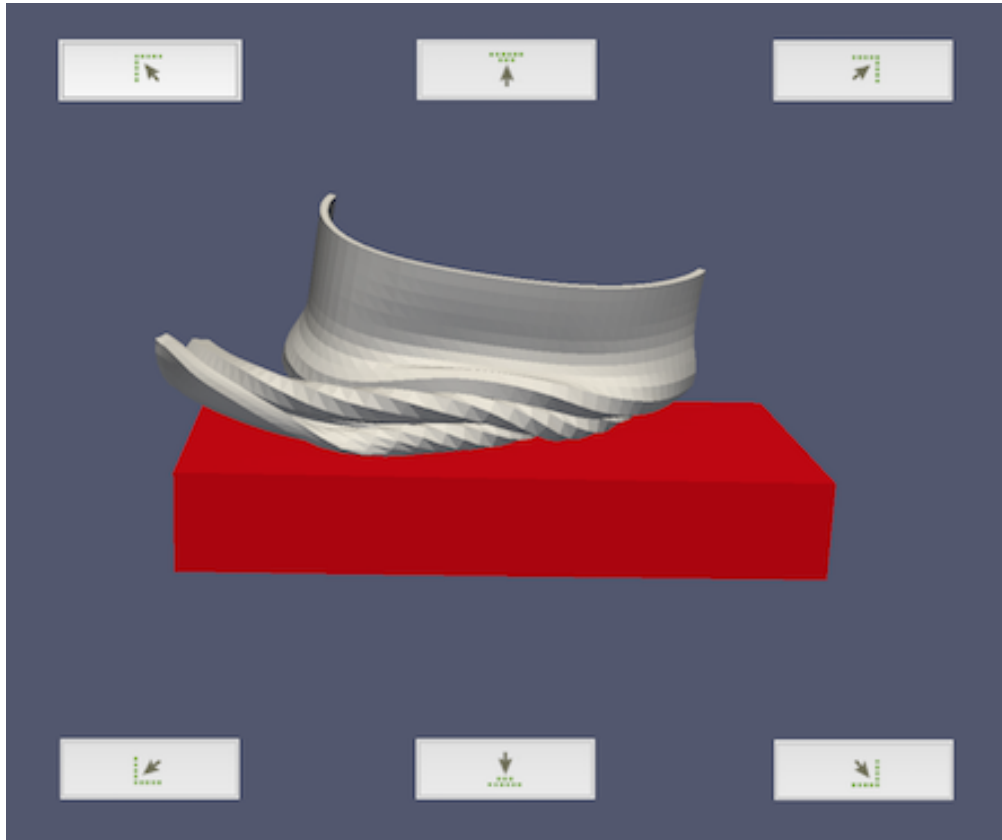


Fig. 2.35: Annotation placement buttons and where they place the annotation.

The second mode, activated by clicking the `Lower Left Corner` checkbox, lets you arbitrarily place the annotation. If the `Interactivity` property is enabled, you can click and drag the annotation in the render view to place it, or you can manually enter a location where the lower left corner of the annotation's bounding box should be placed. The coordinates are defined in terms of fractional coordinates that range from [0, 1] in the x and y dimensions. The coordinate system of the render view has a lower left origin, so a `Lower Left Corner` value of [0, 0] will place the annotation in the lower left corner of the render view.

Text source

The `Text source` enables you to add a text annotation in the render view. It has one property defining what text is displayed. Text can be multiline, and it can contain numbers and unicode characters. Text may also contain Mathtex expressions between starting and ending dollar signs. Mathtext expressions are a subset of TeX math expressions [dt]. When Mathtext is used, the text can only be on a single line.

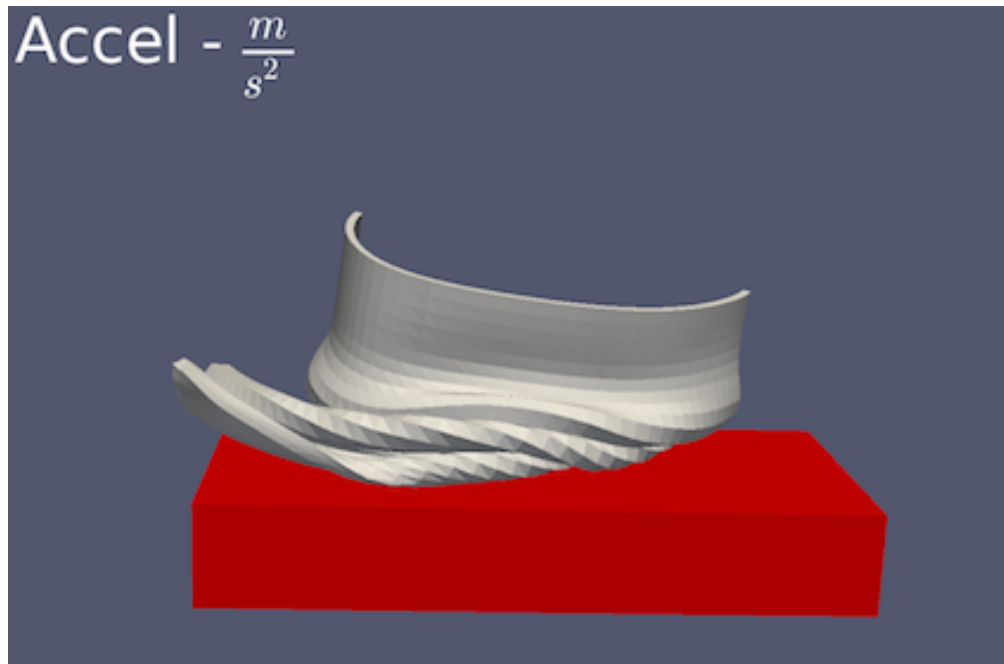


Fig. 2.36: An example of the `Text` source annotation in the upper left corner with a math expression rendered from a Mathtext [dt] expression.

Annotate Time source

The `Annotate Time` source is nearly identical to the `Text` source, but it also offers access to the current time value set in ParaView. The time value is placed in the first format specifier in the `Format` text property. The format specifier can be any number format valid for `printf` and related functions in the C standard library. The default format results in rendering the time value with six decimal digits of precision. Only one format specifier is valid in the property. Additional specifiers after the first will produce undefined results in the annotation text.

2.9.2 Annotation filters

The annotation sources described in the previous section are available for adding text annotations that do not depend on any loaded datasets. To create annotations that show values from an available data source in the `Pipeline Browser`, several annotation filters are available. The properties available to change the text font and annotation location are exactly the same as those available for the annotation sources described in the previous section.

Annotate Attribute Data filter

The `Annotate Attribute Data` makes it possible to create an annotation with a data value from an array (or attribute) in a dataset. To use the filter, first select the data array with the data of interest in the `Select Input Array`. These arrays may be point, cell, or field data arrays. The `Element Id` property specifies the index of the point or cell whose value should be shown in the annotation. If the selected input array is a field array (not associated with points or cells), the `Element Id` specifies the tuple of the array to show. When running in parallel, the `Process Id` denotes the process that holds the array from which the value should be obtained.

The `Prefix` text property precedes the attribute value in the rendered annotation. There is no formatting string - the number is appended after the prefix. If the array value selected is a scalar value, the annotation will contain just the number. On the other hand, if the array value is from a multicomponent array, the individual components will be added to the annotation label in a space-separated list that is surrounded by parentheses.

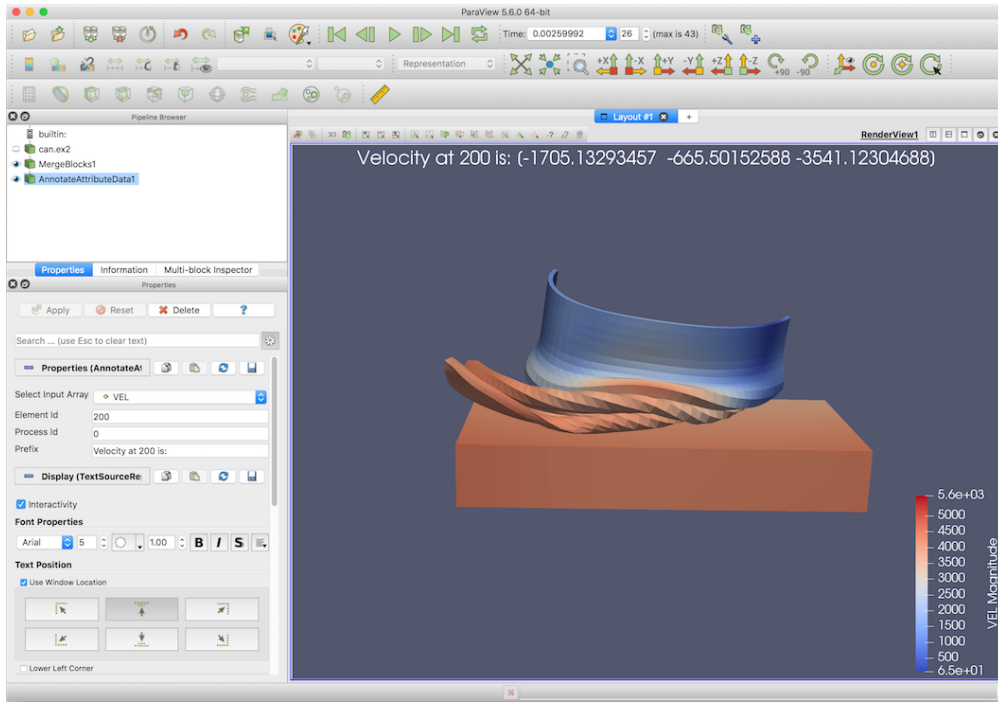


Fig. 2.37: Properties of the `Annotate Attribute Data` filter.

Annotate Global Data filter

Some file formats include the concept of `global` data, a single data value stored in the data array for each time step. **ParaView** stores the set of such data values as a field data array associated with the dataset with the same number of values as timesteps. To display these global values in the render view, use the `Annotate Global Data` filter. The `Select Arrays` popup menu shows the available field data arrays. The `Prefix` and `Suffix` properties come before and after the data value in the annotation, respectively. The `Format` property is a C language number format specifier.

Annotate Time Filter

A nice feature of **ParaView** is that it supports data sources that produce different data at different times. Examples include file readers that read in data for a requested time step and certain temporal filters. Each data source advertises to **ParaView** the time values for which it can produce data. The data produced and displayed in ParaView depends on the time you set in the **ParaView** VCR Controls or Time Inspector panel.

What is even nicer is that you can have several data sources that each advertise and respond to a possibly unique set of times. That is, available sources do not need to advertise that they support the same set of time points - in fact, they may define data at entirely different time points. Given a requested time, each data source will produce the data corresponding to the time it supports closest to the requested time. This features makes it possible to create animations from multiple datasets varying at different time resolutions, for instance.

While the `Annotate Time` source can be used to display **ParaView**'s currently requested time, it does not show the time value to which a particular data source is responding. For example **ParaView** may be requesting data for time 5.0, but if a source produces data for time values 10.0 and above, it will produce the data for time 10.0, even though time 5.0 was requested. To show the time for which a data source is producing data, you can use the `Annotate Time Filter`. Simply attach it to the source of interest. If several data sources are present, a separate instance of this filter may be attached to each one.

This property includes `Shift` and `Scale` parameters used to linearly transform the displayed time. The time value is

first multiplied by the scale and then the shift is then added to it.

Environment Annotation filter

If you want to display information about the environment in which a visualization was generated, use the Environment Annotation filter. By attaching this filter to a data source, you can have it automatically display your user name on the system running ParaView, show which operating system was used to generate it, present the date and time when the visualization was generated, and show the file name of the source data if applicable. Each of these items can be enabled or disabled by checkboxes in the Properties panel for this filter.

If the input source for this filter is a file reader, the File Name property is initialized to the name of the file. A checkbox labeled Display Full Path is available to show the full path of the file, but if unchecked, only the file name will be displayed. This default file path can be overridden by changing the text in the File Name property. If this filter is attached to a filter instead of a reader, the file path will be initialized to an empty string. It can be changed to the original file name manually, or an arbitrary string if so desired.

Python Annotation filter

The most versatile annotation filter, the Python Annotation filter, offers the most general way of generating annotations that include information about the dataset. Values from point, cell, field, and row data arrays may be accessed and combined with mathematical operations in a short Python expression defined in the Expression property. The type of data arrays available for use in the Expression is set with the Array Association property.

Before going further, let's look at an example of how to use the Python Annotation filter. Assume you want to show a data value at from a point array named Pressure at point index 22. First, set the Array Association to Point Data to ensure point data arrays can be referenced in the Python annotation expression. To show the pressure value at point 22, set the Expression property to

Pressure[22]

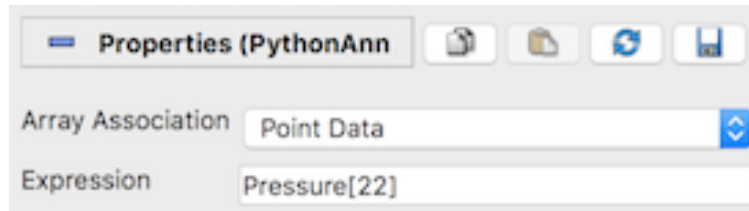


Fig. 2.38: An example of a basic Python Annotation filter showing the value of the Pressure array at point 22.

You can augment the Python expression to give the annotation more meaning. To add a prefix, set the Expression to

'Pressure: %f' % (Pressure[22])

All data arrays in the chosen association are provided as variables that can be referenced in the expression as long as their names are valid Python variables. Array names that are invalid Python variable names are available through a modified version of the array name. This sanitized version of the array name consists of the subset of characters in the array name that are letters, numbers, or underscore (_) joined together without spaces in the order in which they appear in the original array name. For example, an array named Velocity X will be made available in the variable VelocityX.

Point and cell data in composite datasets such as multiblock datasets is accessed somewhat differently than point or cell data in non-composite datasets. The expression

Pressure[22]

retrieves a single scalar value from a point array in a non-composite dataset, the same expression retrieves the 22nd element of the Pressure array in each block. These values are held in a VTKCompositeDataArray, which is a data structure that holds arrays associated with each block in the dataset. Hence, when the expression

Pressure[22]

is evaluated on a composite dataset, the value returned and displayed is actually an assemblage of array values from each block. To access the value from a single block, the array from that block must be selected from the `Arrays` member of the result VTKCompositeDataArray. To show the Pressure value associated with 22nd point of block 2, for example, set the expression to

Pressure[22].Arrays[2]

This expression yields a single data value in the rendered annotation, assuming that the Pressure array has a single component. To show a range of array values, use a Python range expression in the index into the Pressure field, e.g.,

Pressure[22:24].Arrays[2]

This will show the Pressure values for points 22 and 23 from block 2. You can also retrieve more than one array using an index range on the `Arrays` member, e.g.,

Pressure[22:24].Arrays[2:5]

This expression evaluates to Pressure for points 22 and 23 for blocks 2, 3, and 4.

The `Array Association` is really a convenience to make the set of data arrays of the given association available as variables that can be used in the `Expression`. The downside of using these array names is that arrays from only one array association are available at a time. That means annotations that require the combination of a cell data array and point data array, for example, cannot be expressed with these convenience Python variables alone.

Fortunately, you can access any array in the input to this filter with a slightly more verbose expression. For example, the following expression multiplies a cell data value by a point data value:

```
inputs[0].CellData['Volume'][0] * inputs[0].PointData['Pressure'][0]
```

Note that the arrays in the input are accessed in the above example using their original array names.

In the example above, the expression `inputs[0]` refers to the first input to the filter. While this filter can take only one input, it is based on the same code used by the `Python Calculator` (described in [Section 1.5.9](#)), which puts its several inputs into a Python list, hence the input to the `Python Annotation` filter is referenced as `inputs[0]`.

In addition to making variables for the current array association available in the expression, this filter provides some other variables that can be useful when computing an annotation value.

- `points` : Point locations (available for datasets with explicit points).
- `time_value` , `t_value` : The current time value set in **ParaView**.
- `time_steps` , `t_steps` : The number of timesteps available in the input.
- `time_range` , `t_range` : The range of timesteps in the input.
- `time_index` , `t_index` : The index of the current timestep in **ParaView**.

There are some situations where the variables above are not defined. If the input has no explicitly defined points, e.g., image data, the `points` variable is not defined. If the input does not define timesteps, the `time_*` and `t_*` variables are not defined.

Finally, all the capabilities of the Python Calculator, documented in [Section 1.5.9](#), are available, including the NumPy integration and access to the NumPy and SciPy methods.

Common Errors

The time-related variables are not needed to index into point or cell data arrays. Only the point and cell arrays loaded for the current timestep are available in the filter. You cannot access point or cell data from arbitrary timesteps from within this filter.

With the capabilities in this filter, it is possible to reproduce the other annotation sources and filters, as shown below.

- **Text source:** To produce the text “My annotation”, write "My annotation"
- **Annotate Time source:** To produce the equivalent of Time: %f, write "Time :%f" % time_value
- **Annotate Attribute Data filter:** To produce the equivalent of setting Select Input Array to EQPS, Element Id to 0 and Process Id to 0, and Prefix to Value is:, write 'Value is: %.12f' % (inputs[0].CellData['EQPS'][0]) .
- **Annotate Global Data filter:** To produce the same annotation as setting Select Arrays to KE, Prefix to Value is: , Format to %7.5g, and empty suffix, write "Value is: %7.5g" % (inputs[0].FieldData['KE'].Arrays[0][time_index])
- **Annotate Time Filter:** To produce the equivalent of setting Format to Time: %f, Shift to 3, and Scale to 2, write "Time: %f" % (2*time_value + 3) .

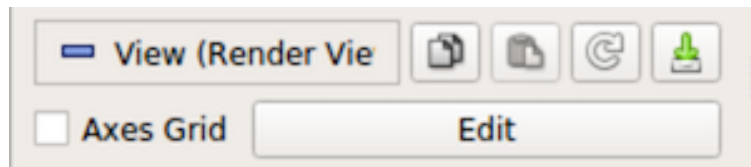
The examples above are meant to illustrate the versatility of the Python Annotation filter. Using the specialized annotation sources and filters are likely to be more convenient than entering the expressions in the examples.

2.10 Axes Grid

Oftentimes, you want to render a reference grid in the background for a visualization – think axes in a chart view, except this time we are talking of the 3D Render View . Such a grid is useful to get an understanding for the data bounds and placement in 3D space. In such cases, you use the Axes Grid . Axes Grid renders a 3D grid with labels around the rendered scene. In this chapter, we will take a closer look at using and customizing the Axes Grid .

2.10.1 The basics

To turn on the Axes Grid for a Render View , you use the Properties panel. Under the View section, you check the Axes Grid checkbox to turn the Axes Grid on for the active view.



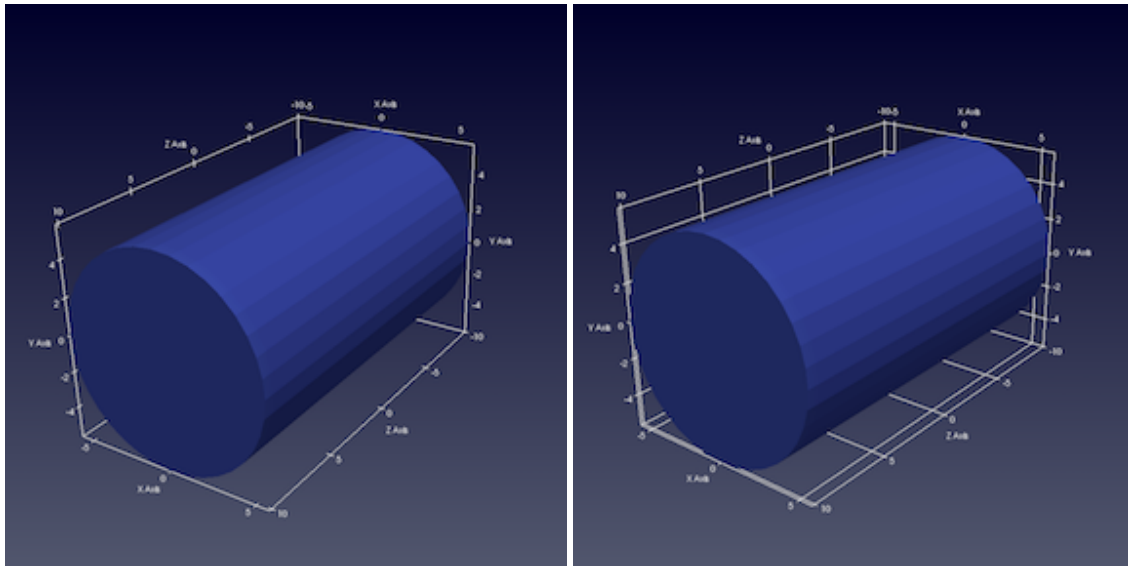
Clicking on the Edit button will pop up the Axes Grid properties dialog ([Fig. 2.39](#)) that allows you to customize the Axes Grid . As with the Properties panel, this is a searchable dialog, hence you can use the Search box at the top of the dialog to search of properties of interest. At the same time, the button can be used to toggle between default and advanced modes for the panel.

Using this dialog, you can change common properties like the titles (X Title , Y Title , and Z Title), title and label fonts using Title Font Properties and Label Font Properties for each of the axes directions, as well as

the `Grid Color`. Besides labelling the axes, you can render a grid by checking `Show Grid`. Once you have the `Axes Grid` setup to your liking, you can use the  to save your selections so that they are automatically loaded next time you launch ParaView. You can always use the  to revert back to ParaView defaults.

2.10.2 Use cases

To get a better look at the available customizations, let's look at various visualizations possible and then see how you can set those up using the properties on the `Edit Axes Grid` dialog. In these examples, we use the `disk_out_ref.ex2` example dataset packaged with ParaView.



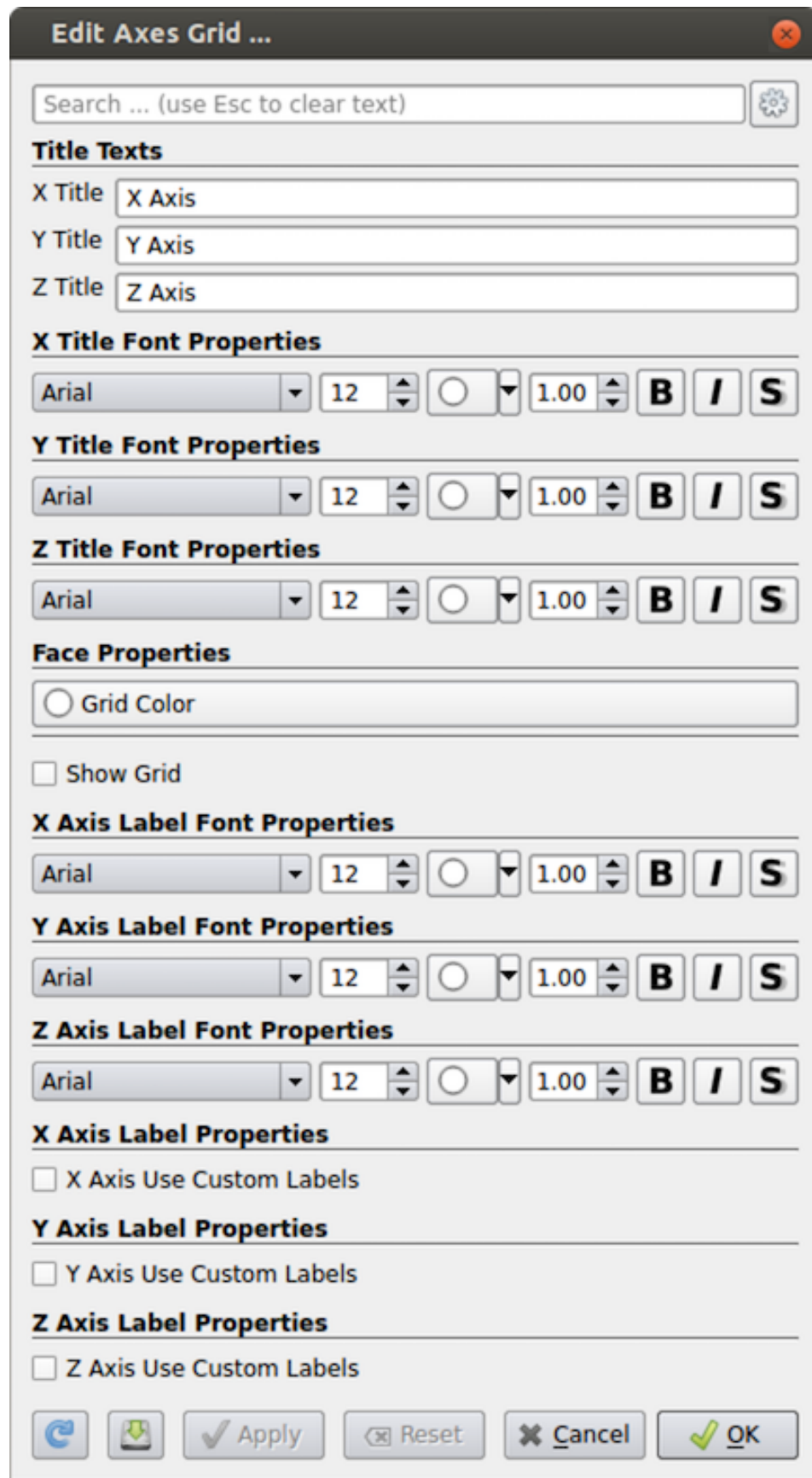
In the images above, on the left is the default `Axes Grid`. Simply turning on the visibility of the `Axes Grid` will generate such a visualization. The axes places always stay behind the rendered geometry even as you interact with the scene. As you zoom in and out, the labels and ticks will be updated based on visual cues.

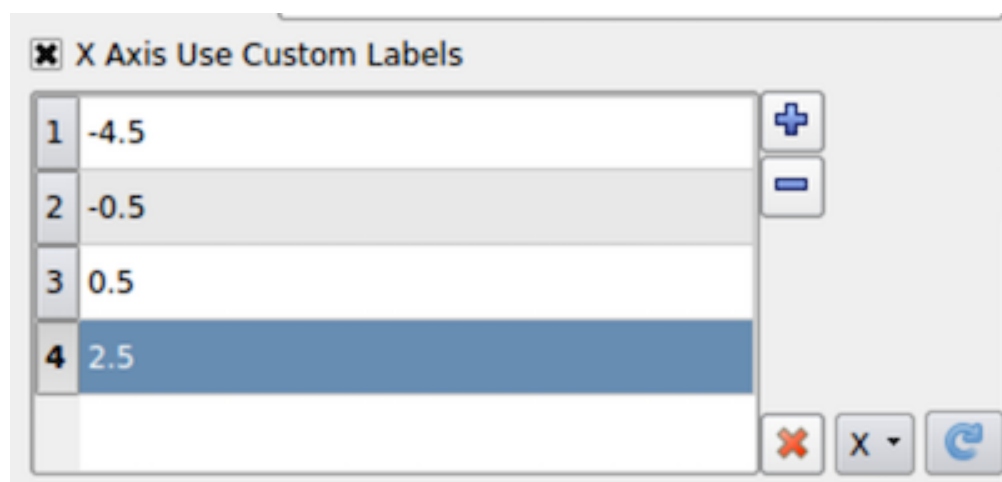
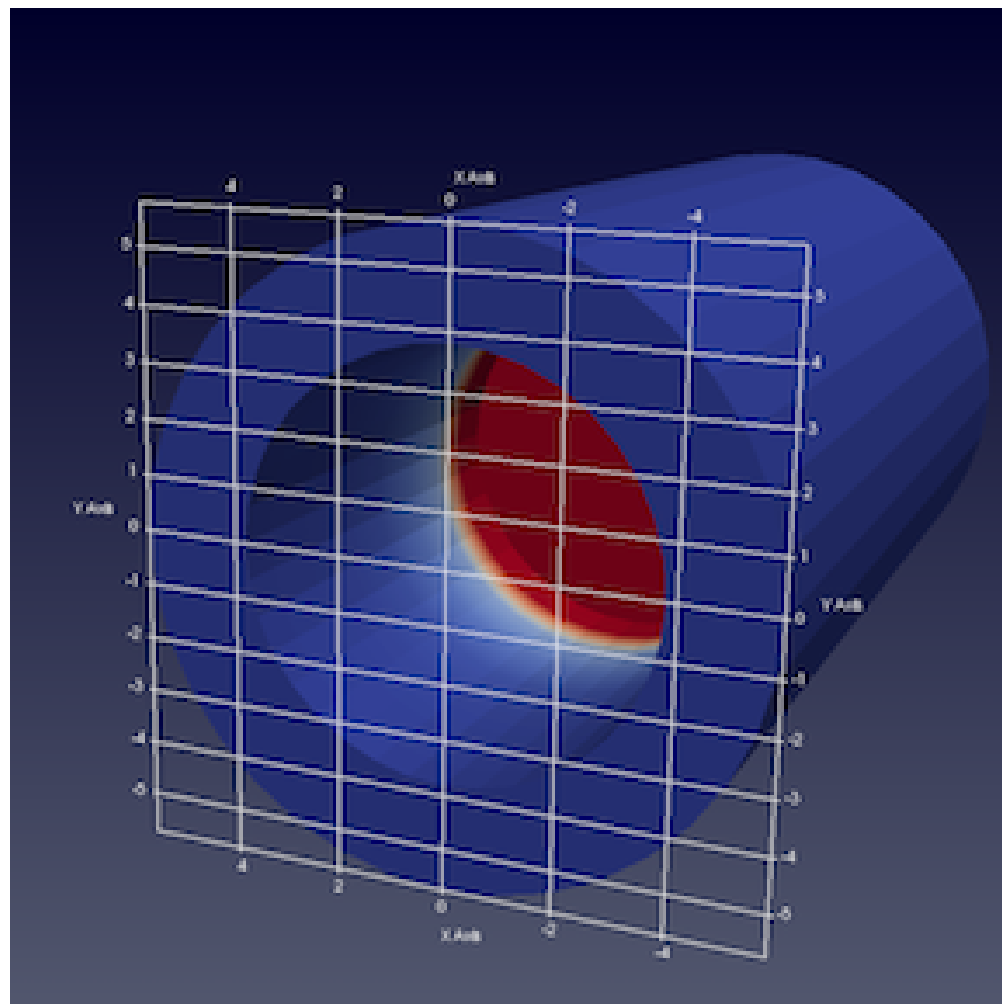
To show a grid along the axes planes, aligned with the ticks and labels, turn on the `Show Grid` checkbox, resulting in a visualization on the right.

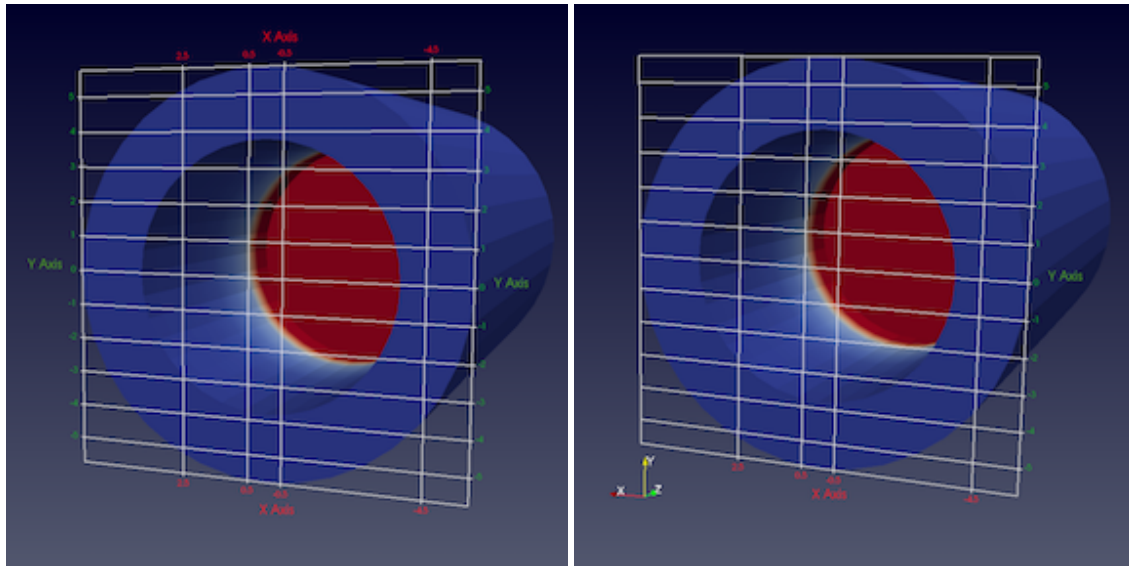
By default, the gridded faces are always the farthest faces i.e. they stay behind the rendered geometry and keep on updating as you rotate the scene. To fix which faces of the bounding-box are to be rendered, use the `Faces To Render` button (it's an advanced property, so you may have to search for it using the Search box in the `Edit Axes Grid` dialog). Suppose, we want to label just one face, the lower XY face. In that case, uncheck all the other faces except `Min-XY` in menu popped up on clicking on the `Faces to Render` button. This will indeed just show the min-XY face, however as you rotate the scene, the face will get hidden as soon as the face gets closer to the camera than the dataset. This is because, by default, `Cull Frontfaces` is enabled. Uncheck `Cull Frontfaces` and ParaView will stop removing the face as it comes ahead of the geometry, enabling a visualization as follows.

Besides controlling which faces to render, you can also control where the labels are placed. Let's say we want ParaView to decide how to place labels along the Y axis, however for the X axis, we want to explicitly label the values 2.5, 0.5, -0.5 , and -4.5 . To that, assuming we are the advanced mode for the `Edit Axes Grid` panel, check `X Axis Use Custom Labels`. That will show a table widget that allows you to add values as shown below.

Using the  button, add the custom values. While at it, let's also change the `X Axis Label Font Properties` and `X Axis Title Font Properties` to change the color to red and similar for the Y axis, let's change the color to green. Increase the title font sizes to 18, to make them stand out and you will get a visualization as follows (below, left).







Here we see that both sides of the axis plane are labeled. Suppose you only want to label one of the sides, in that case use the `Axes To Label` property to uncheck all but `Min-X` and `Min-Y`. This will result in the visualization shown above, right.

2.10.3 Axes Grid in pvython

In pvython, `Axes Grid` is accessible as the `AxesGrid` property on the render view.

```
>>> renderView = GetActiveView()

# AxesGrid property provides access to the AxesGrid object.
>>> axesGrid = renderView.AxesGrid

# To toggle visibility of the axes grid,
>>> axesGrid.Visibility = 1
```

All properties on the `Axes Grid` that you set using the `Edit Axes Grid` dialog are available on this `axesGrid` object and can be changed as follows:

```
>>> axesGrid.XTitle = 'X Title'
>>> axesGrid.XTitleColor = [0.6, 0.6, 0.0]
>>> axesGrid.XAxisLabels = [-0.5, 0.5, 2.5, 3.5]
```

Note you can indeed use the tracing capabilities described in [Section 1.1.6](#) to determine what Python API to use to change a specific property on the `Edit Axes Grid` dialog or use `help`.

```
>>> help(axesGrid)
Help on GridAxes3DActor in module paraview.servermanager object:

class GridAxes3DActor(Proxy)
|   GridAxes3DActor can be used to render a grid in a render view.
|
|   Method resolution order:
|       GridAxes3DActor
|       Proxy
```

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```

| __builtin__.object

| Methods defined here:

|     Initialize = aInitialize(self, connection=None, update=True)

|     -----
|     Data descriptors defined here:

|         AxesToLabel
|             Set the mask to select the axes to label. The axes labelled will be a subset of the
|             axes selected depending on which faces are also being rendered.

|         CullBackface
|             Set to true to hide faces of the grid facing away from the camera i.e. hide all
|             back faces.

|         CullFrontface
|             Set to true to hide faces of the grid facing towards from the camera i.e. hide all
|             front faces.

|         DataPosition
|             If data is being translated, you can show the original data bounds for the axes
|             instead of the translated bounds by setting the DataPosition to match the
|             translation applied to the dataset.

|     ...

```

2.11 Customizing ParaView

ParaView can be customized in a number of ways to tailor it to your preferences and needs. Customization options include setting general application behavior, customizing default property values used for filters, representations, and views, and customizing aspects of the paraview client. This chapter describes the different ways to customize **Paraview**.

2.11.1 Settings

As with any large application, paraview provides mechanisms to customize some of its application behavior. These are referred to as **application settings** or just **settings**. Such settings can be changed using the **Settings** dialog, which is accessed from the *Edit > Settings* menu (*ParaView > Preferences* on the Mac). We have seen parts of this dialog earlier, e.g., in [Section 2.1.2](#), [Section 2.6.12](#), and [Section 2.6.12](#). In this section, we will take a closer look at some of the other options available in this dialog.

The **Settings** dialog is split into several tabs. The **General** tab consolidates most of the miscellaneous settings. The **Camera** tab enables you to change the mouse interaction mappings for the **Render View** and similar views. The **Render View** tab, which we saw earlier in [Section 2.6.12](#) and [Section 2.6.12](#), provides options in regards to rendering in **Render View** and similar views. The **Color Palette** tab is used to change the active color palette.

Using this dialog is not much different than the **Properties** panel. You have the **Search** box at the top, which allows you to search properties matching the input text ([Section 2.1.1](#)). The  button can be used to toggle between default and advanced modes.

To apply the changes made to any of the settings, use the **Apply** or **OK** buttons. **OK** will apply the changes and close the dialog, while **Cancel** will reject any changes made and close the dialog. Any changes made to the options in this dialog are persistent across sessions. That is, the next time you launch paraview, you'll still be using the same settings chosen earlier. To revert to the default, use the **Restore Defaults** button. You can also manually edit the setting file as in [Section 2.11.2](#). Furthermore, site maintainers can provide site-wide defaults for these, as is explained in [Section 2.11.2](#).

Next, we will see some of the important options available. Those that are only available in the advanced mode are indicated as such using the  icon. You will either need to toggle on the advanced options with the  button or search for the option using the **Search** box.

General settings

- General Options
 - **Show Welcome Dialog** : Uncheck this to not show the welcome screen at application startup. You will need to restart paraview to see the effect. 
 - **Show Save State On Exit** : When this is checked paraview will prompt you to save a state file when you exit the application. 
 - **Crash Recovery** : When this is checked, paraview will intermittently save a backup state file as you make changes in the visualization pipeline. If paraview crashes for some reason, then when you relaunch paraview, it will provide you with a choice to load the backup state saved before the crash occurred. This is not 100% reliable, but some users may find it useful to avoid losing their visualization state due to a crash. 
 - **Force Single Column Menus** : On platforms that support multicolumn menus, ensure all menu items are selectable on low-resolution screens.
- GUI Font
 - **Override Font** : When checked, use a custom font size for the user interface. This overrides the system default font size.
 - **Font Size** : The size of the font to use for UI elements.
- View Options
 - **Default View Type** : When paraview starts up, it creates **Render View** by default. You can use this option to change the type of the view that is created by default, instead of the **Render View**. You can even pick **None** if you don't want to create any view by default. 
- Properties Panel Options
 - **Auto Apply** : When checked, the **Properties** panel will automatically apply any changes you make to the properties without requiring you to click the **Apply** button. The same setting can also be toggled using the  button in the **Main Controls** toolbar.
 - **Auto Apply Active Only** : This limits the auto-applying to the properties on the active source alone. 

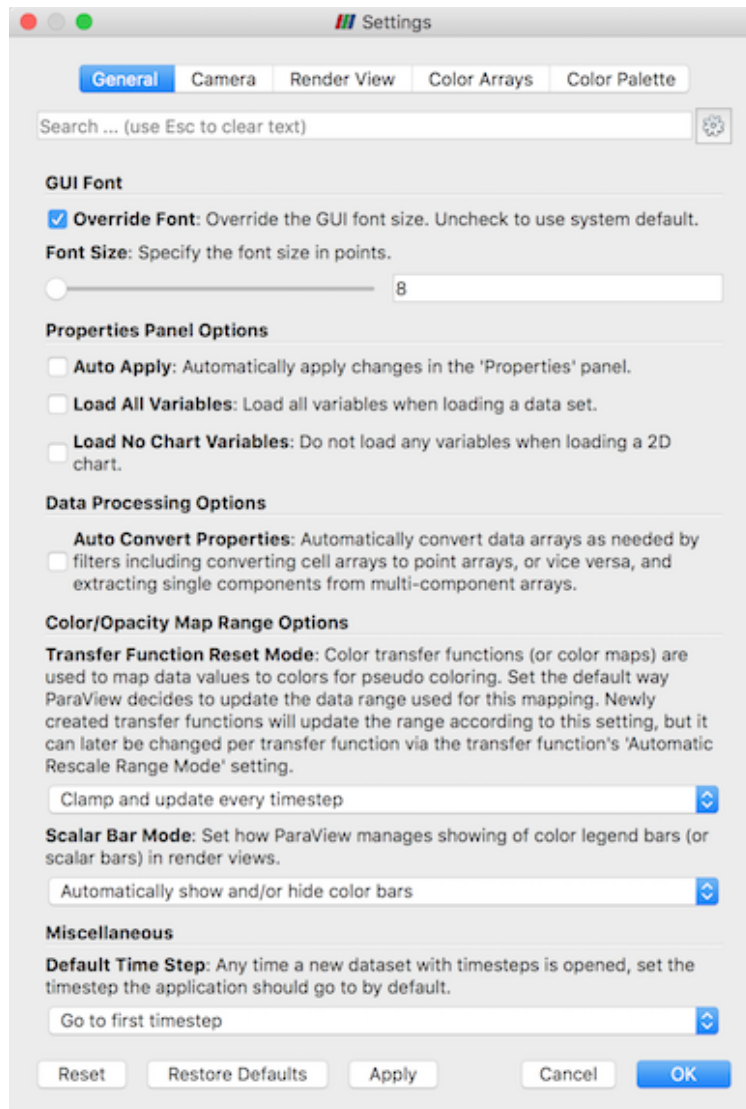


Fig. 2.40: Settings dialog in paraview showing the General settings tab.

- **Properties Panel Mode** : This allows you to split the Properties panel into separate panels as described in [Section 2.1.2](#). 
- **Data Processing Options**
 - **Auto Convert Properties** : Several filters only work on one type of array, e.g., point data arrays or cell data arrays. Before using such filters, you are expected to apply the Point Data To Cell Data or Cell Data To Point Data filters. To avoid having to add these filters explicitly, you can check this checkbox. When checked, **ParaView** will automatically convert data arrays as needed by filters, including converting cell array to point arrays and vice-versa, as well as extracting a single component from a multi-component array.
- **Color/Opacity Map Range Options**
 - **Transfer Function Reset Mode** : This setting controls the initial settings for how **ParaView** will reset the ranges for color and opacity maps (or transfer functions). This sets the initial value of the Automatic Rescale Range Mode for newly created color-opacity maps ([Section 2.2.2](#)). This setting can be changed on a per-color map basis after the color map has been created.
 - **Scalar Bar Mode** : This setting controls how paraview manages showing the color legend (or scalar bar) in Render View and similar views.
- **Default Time Step** Whenever a dataset with timesteps is opened, this setting controls how paraview will update the current time shown by the application. You can choose between Leave current time unchanged, if possible, Go to first timestep, and Go to last timestep.
- **Animation**
 - **Cache Geometry For Animation** : This enables caching of geometry when playing animations to attempt to speed up animation playback in a loop. When caching is enabled, data ranges reported by the Information panel and others can be incorrect, since the pipeline may not have updated. 
 - **Animation Geometry Cache Limit** : When animation caching is enabled, this setting controls how much geometry (in kilobytes) can be cached by any rank. As soon as a rank's cache size reaches this limit, **ParaView** will no longer cache the remaining timesteps. 
 - **Animation Time Notation** : Sets the display notation for the time in the annotation toolbar. Options are Mixed, Scientific, and Fixed .
 - **Animation Time Precision** : Sets the number of digits displayed in the time in the animation toolbar.
- **Maximum Number of Data Representation Labels** When a selection is labeled by data attributes this is the maximum number of labels to use. When the number of points/cells to label is above this value then a subset of this many will be labeled instead. Too many overlapping labels becomes illegible, so this is set to 100 by default.

Camera settings

This tab allows you to control how you can interact in Render View and similar views. Basically, you are setting up a mapping between each of the mouse buttons and keyboard modifiers, and the available interaction types including Rotate, Pan, Zoom, etc. The dialog allows you to set the interaction mapping separately for 3D and 2D interaction modes (see [Section 1.4.4](#)).

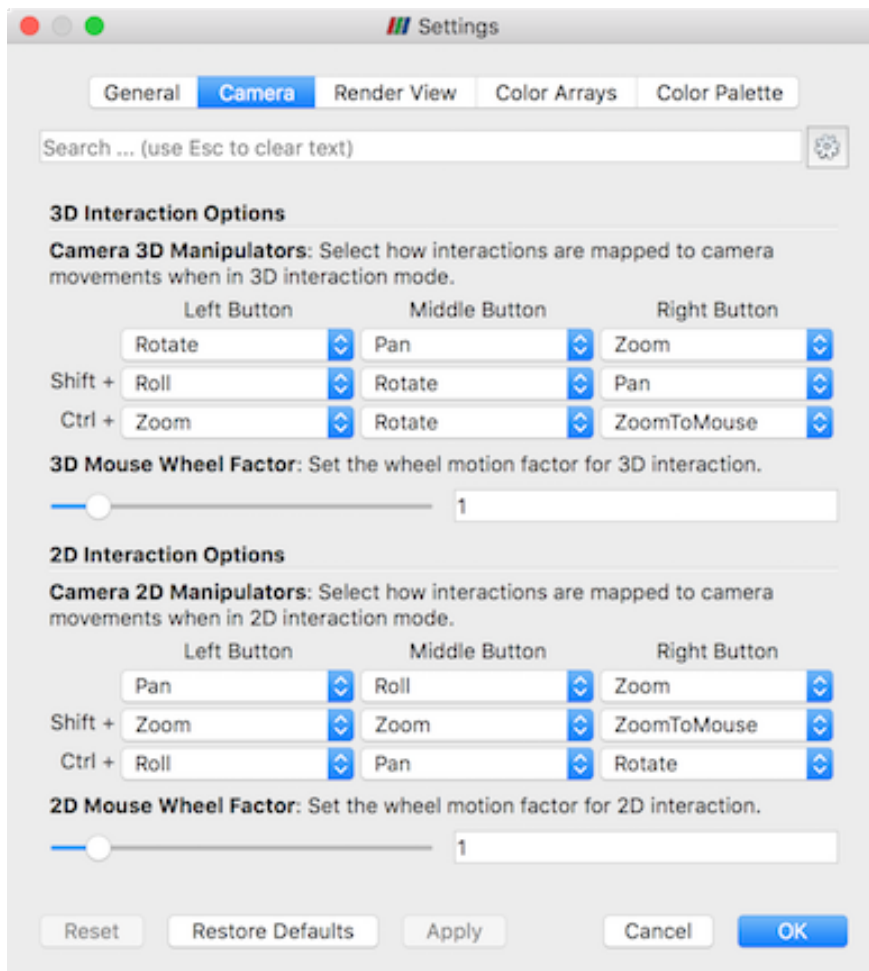


Fig. 2.41: Settings dialog in paraview showing the Camera settings tab.

Render View settings

Refer to [Section 2.6.12](#) and [Section 2.6.12](#) for various options available on the Render View tab.

Color Palette

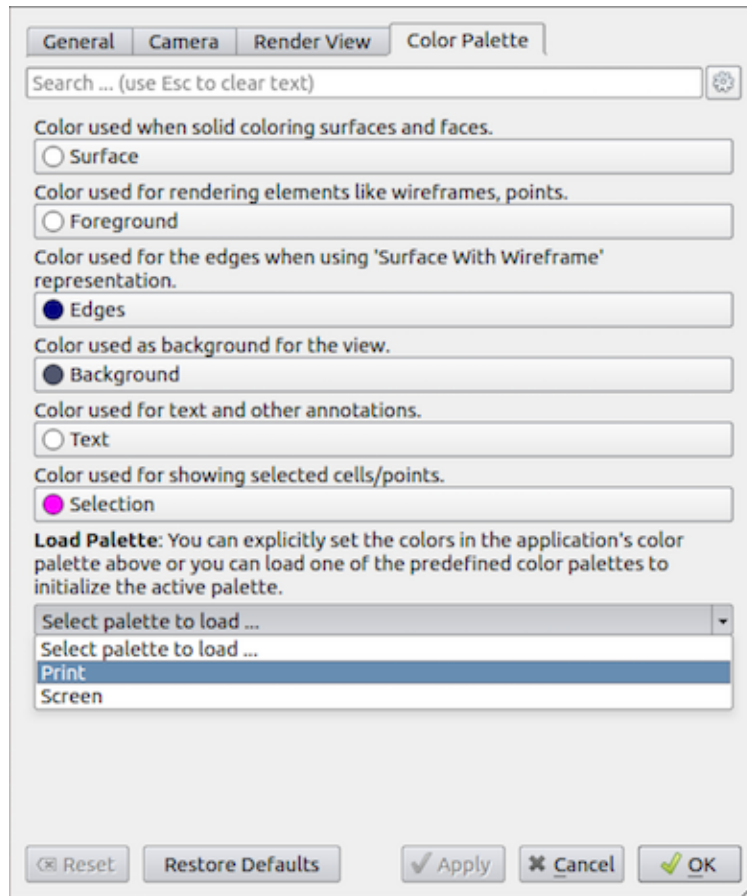


Fig. 2.42: Settings dialog in paraview showing the Color Palette settings tab.

The Color Palette tab ([Fig. 2.42](#)) allows you to change the colors in the active color palette. The tab lists the available color categories `Surface`, `Foreground`, `Edges`, `Background`, `Text`, and `Selection`. You can manually set colors to use for each of these categories or load one of the predefined palettes using the `Load Palette` option. To understand `color palettes`, let's look at an example.

Let's start `paraview` and split the active view to create two `Render View` instances side by side. You may want to start `paraview` with the `-dr` command line argument to stop any of your current settings from interfering with this demo. Next, show `Sphere` as `Wireframe` in the view on the left, and show `Cone` as `Surface` in the view on the right. Also, turn on `Cube Axis` for `Cone`. You will see something like [Fig. 2.43](#) (top).

Now let's say you want to generate an image for printing. Typically, for printing, you'd want the background color to be white, while the wireframes and annotations to be colored black. To do that, one way is to go change each of the colors for each each of the views, displays and cube-axes. You can imagine how tedious that will get especially with larger pipelines. Alternatively, using the `Settings` dialog, change the active color palette to `Print` as shown in [Fig. 2.42](#) and then click `OK` or `Apply`. The visualization will immediately change to something like [Fig. 2.43](#) (bottom).

Essentially, **ParaView** allows you to *link* any color property to one of the color categories. When the color palette is changed, any color property linked to a palette category will also be automatically updated to match the category

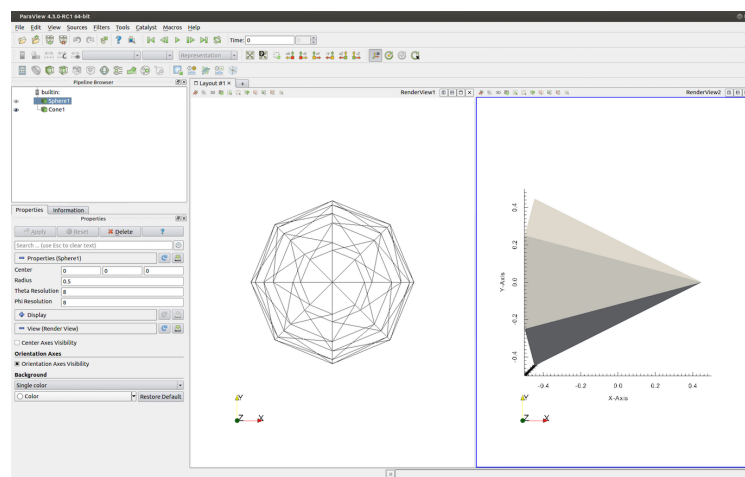
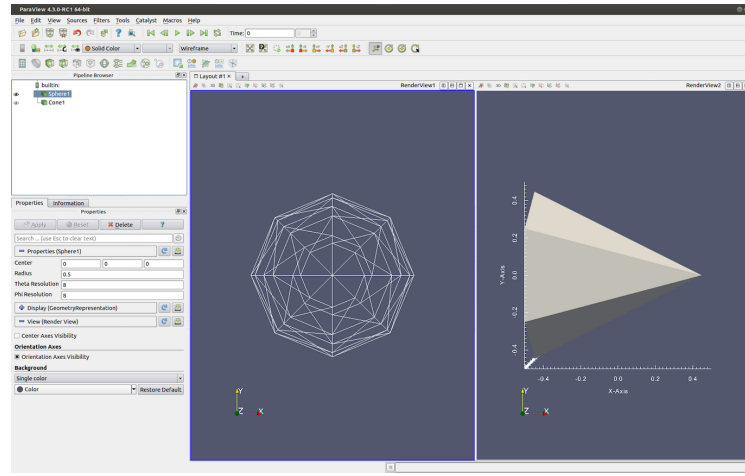


Fig. 2.43: The effect of loading the Print color palette as the active palette. The top is the original visualization and the bottom shows the result after loading the Print palette.

color. Fig. 2.44 shows how to link a color property to a color palette category in the Properties panel. Use the tiny drop-down menu marker to make the menu pop up that shows the color palette categories. Select any one of them to link that property with the category. The link is automatically severed if you manually change the color by simply clicking on the button.

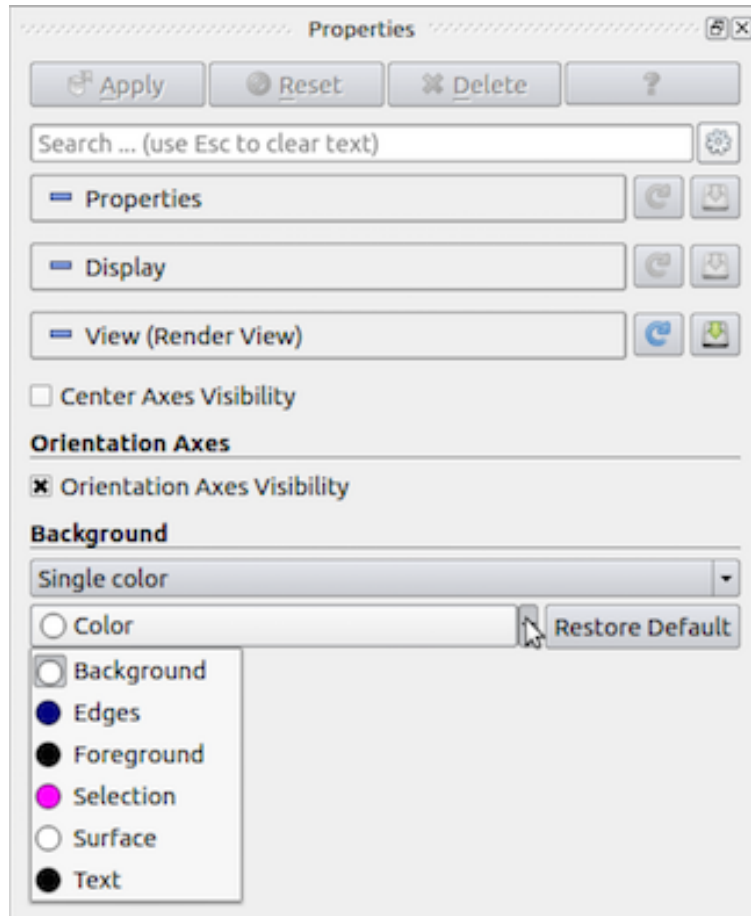


Fig. 2.44: Popup menu allows you to link a color property to a color palette category in the Properties panel.}

2.11.2 Custom default settings

The section describes how to specify custom default settings for the properties of sources, readers, filters, representations, and views. This can be used to specify, for example, the default background color for new views, whether a gradient background should be used, the resolution of a sphere source, which data arrays to load from a particular file format, and the default setting for almost any other object property.

The same custom defaults are used across all the **ParaView** executables. This means that custom defaults specified in the `paraview` executable are also used as defaults in `pypython` and `pvbatch`, which makes it easier to set up a visualization with `paraview` and use `pypython` or `pvbatch` to generate an animation from time-series data, for example.

Customizing defaults for properties

The Properties panel in **paraview** has three sections, **Properties**, **Display**, and **View**. Each section has two buttons. These buttons are circled in red in Fig. 2.45. The button with the disk icon is used to save the current property values in that section that have been applied with the **Apply** button. Property values that have been changed but not applied with the **Apply** button will not be saved as custom default settings.

The button with the circular arrow (or reload icon) is used to restore any custom property settings for the object to **ParaView**'s application defaults. Once you save the current property settings as defaults, those values will be treated as the defaults from then on until you change them to another value or reset them. The saved defaults are written to a configuration file so that they are available when you close and launch **ParaView** again.

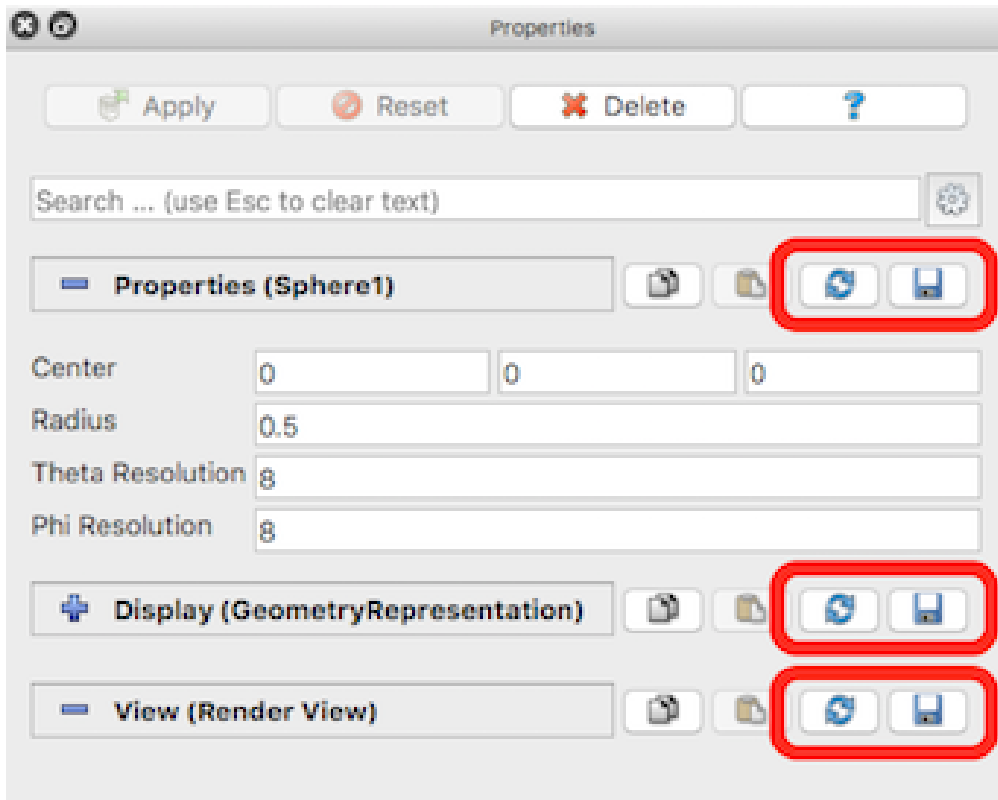


Fig. 2.45: Buttons for saving and restoring default property values in the Properties panel.

You can undo your changes to the default property values by clicking on the reload button. This will reset the current view property values to **paraview**'s application defaults. To fully restore **paraview**'s default values, you need to click the save button again. If you don't, the restored default values will be applied only to the current object, and new instances of that object will have the custom default values that were saved the last time you clicked the save button.

Example: specifying a custom background color

Suppose you want to change the default background color in the Render View . To do this, scroll down to the View section of the Properties panel and click on the combo box that shows the current background color. Select a new color, and click OK . Next, scroll up to the View (Render View) section header and click on the disk button to the right of the header. This will save the new background color as the default background color for new views. To see this, click on the + sign next to the tab above the 3D view to create a new layout. Click on the Render View button. A new render view will be created with the custom background color you just saved as default.

Configuring default settings with JSON

Custom default settings are stored in a text file in the JSON (JavaScript Object Notation) format. We recommend to use the user interface in paraview to set most default values, but it is possible to set them by editing the JSON settings file directly. It is always a good idea to make a backup copy of a settings file prior to manual editing.

The **ParaView** executables read from and write to a file named `ParaView-UserSettings.json`, which is located in your home directory on your computer. On Windows, this file is located at `%APPDATA%/ParaView/ParaView-UserSettings.json`, where the APPDATA environment variable is usually something like `C:/Users/USERNAME/AppData/Roaming`, where USERNAME is your login name. On Unix-like systems, it is located under `~/config/ParaView/ParaView-UserSettings.json`. This file will exist if you have made any default settings changes through the user interface in the paraview executable. Once set, these default settings will be available in subsequent versions of **ParaView**.

A simple example of a file that specifies custom default settings is shown below:

```
{  
  "sources" : {  
    "SphereSource" : {  
      "Radius" : 3.5,  
      "ThetaResolution" : 32  
    },  
    "CylinderSource" : {  
      "Radius" : 2  
    }  
  },  
  "views" : {  
    "RenderView" : {  
      "Background" : [0.0, 0.0, 0.0]  
    }  
  }  
}
```

Note the hierarchical organization of the file. The first level of the hierarchy specifies the group to which the object whose settings are being specified refers (“sources” in this example). The second level names the object whose settings are being specified. Finally, the third level specifies the custom default settings themselves. Note that default values can be set to literal numbers, strings, or arrays (denoted by comma-separated literals in square brackets).

The names of groups and objects come from the XML proxy definition files in **ParaView**’s source code in the directory `ParaView/ParaViewCore/ServerManager/SApplication/Resources` (`ParaView\ParaViewCore\ServerManager\SApplication\Resources` on Windows systems) . The group name is defined by the `name` attribute in a `ProxyGroup` element. The object name comes from the `name` attribute in the `Proxy` element (or elements of `vtkSMProxy` subclasses). The property names come from the `name` attribute in the `*Property` XML elements for the object.

Did you know?

The application-wide settings available in paraview through the *Edit > Settings* menu are also saved to this user settings file. Hence, if you have changed the application settings, you will see some entries under a group named “settings”.

Configuring site-wide default settings

In addition to individual custom default settings, **ParaView** offers a way to specify site-wide custom default settings for a **ParaView** installation. These site-wide custom defaults must be defined in a JSON file with the same structure as the user settings file. In fact, one way to create a site settings file is to set the custom defaults desired in paraview, close the program, and then copy the user settings file to the site settings file. The site settings file must be named `ParaView-SiteSettings.json`.

The **ParaView** executables will search for the site settings file in several locations. If you installed **ParaView** in the directory `INSTALL`, then the **ParaView** executables will search for the site settings file in these directories in the specified order:

- `INSTALL/share/paraview-X.Y` (`INSTALLshareparaview-X.Y` in Windows systems)
- `INSTALL/lib` (`INSTALLlib` in Windows systems)
- `INSTALL`
- `INSTALL/..`. (`INSTALL/lib` in Windows systems)

where `X` is **ParaView**’s major version number and `Y` is the minor version number. **ParaView** executables will search these directories in the given order, reading in the first `ParaView-SiteSettings.json` file it finds. The conventional location for this kind of configuration file is in the `share` directory (the first directory searched), so we recommend placing the site settings file there.

Custom defaults in the user settings file take precedence over custom defaults in the site settings. If the same default is specified in both the `ParaView-SiteSettings.json` file and `ParaView-UserSettings.json` file in a user’s directory, the default specified in the `ParaView-UserSettings.json` file will be used. This is true for both object property settings and application-settings set through the *Edit > Settings* menu.

To aid in debugging problems with the site settings file location, you can define an environment variable named `PV_SETTINGS_DEBUG` to something other than an empty string. This will turn on verbose output showing where the **ParaView** executables are looking for the site settings file.

**CHAPTER
THREE**

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